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(54) Title: PREVENTION OF Bt RESISTANCE DEVELOPMENT

#### (57) £bstract

Plants made resistant to insects by transforming their nuclear genome with two or more DNA sequences, each encoding a different non-competitively binding *B. thuringiensis* protoxin or insecticidal part thereof, preferably the toxin thereof.

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### PREVENTION OF Bt RESISTANCE DEVELOPMENT

This invention relates to plant cells and plants, the genomes of which are transformed to contain at least two genes, each coding for a different noncompetitively binding Bacillus thuringiensis ("B.thuringiensis" or "Bt") insecticidal protein ("ICP") for a specific target insect species, preferably belonging to the order of Lepidoptera or Coleoptera. Such transformed plants have advantages over plants transformed with a single B. thuringiensis ICP gene, especially with respect to the prevention of resistance development in the target insect species against the at least two B. thuringiensis expressed in such plants.

This invention also relates to a process for the production of such transgenic plants, taking into account the competitive and non-competitive binding properties of the at least two B. thuringiensis ICPs in the target insect species' midgut. Simultaneous expression in plants of the at least two genes, each coding for a different non-competitively binding B. thuringiensis ICP in plants, is particularly useful to prevent or delay resistance development of insects against the at least two B. thuringiensis ICPs expressed in the plants.

This invention further relates to a process for the construction of novel plant expression vectors and to the novel plant expression vectors themselves, which contain the at least two <u>B. thuringiensis</u> ICP genes encoding the at least two non-competitively binding <u>B. thuringiensis</u> ICPs. Such vectors allow integration and coordinate expression of the at least two <u>B. thuringiensis</u> ICP genes in plants.

### BACKGROUND OF THE INVENTION

Since the development and the widespread use of chemical insecticides, the occurrence of resistant important problem. strains has been an insect Development of insecticide resistance is a phenomenon dependent on biochemical, physiological, genetic and ecological mechanisms. Currently, insect resistance has been reported against all major classes of chemical including chlorinated hydrocarbons, organophosphates, carbamates, and pyrethroid compounds (Brattsten et al., 1986).

In contrast to the rapid development of insect resistance to synthetic insecticides, development of insect resistance to bacterial insecticides such as B. thuringiensis sprays has evolved slowly despite many years of use (Brattsten et al., 1986). The spore forming gram-positive bacterium В. thuringiensis produces a parasporal crystal which is composed of crystal proteins (ICPs) having insecticidal activity. Important factors decreasing the probability emergence of resistant insect strains in the field against B. thuringiensis sprays are: firstly the short half-life of B. thuringiensis sprays after foliar application; secondly the fact that commercial B. thuringiensis preparations often consist of a mixture of several insecticidal factors including spores, ICPs and eventually beta-exotoxins (Shields, 1987); the transitory nature of plant-pest interactions. Many successful field trials have shown that commercial preparations of a B. thuringiensis containing its spore-crystal complex, effectively control lepidopterous pests in agriculture and forestry (Krieg and Langenbruch, 1981). B. thuringiensis is at present the most widely used pathogen for microbial control of insect pests.

PCT/EP90/00905

Various laboratory studies, in which selection against <u>B</u>. <u>thuringiensis</u> was applied over several generations of insects, have confirmed that resistance against <u>B</u>. <u>thuringiensis</u> is seldom obtained. However, it should be emphasized that the laboratory conditions represented rather low selection pressure conditions.

For example, Goldman et al. (1986) have applied selection with B. thuringiensis israelensis toxin over 14 generations of Aedes aegypti and found only a marginal decrease in sensitivity. The lack of any observable trend toward decreasing susceptibility in the selected strains may be a reflection of the low selection pressure (LC50) carried out over a limited number of generations. However, it should be pointed out that Georghiou et al. (In : Insecticide Resistance in Mosquitoes : Research on new chemicals techniques "Mosquito Control for management. In Research. Annual Report 1983, University California.") with <u>Culex quinquefasciatus</u> obtained an 11-fold increase in resistance to B. thuringiensis israelensis after 32 generations at LC<sub>95</sub> selection presssure.

McGaughey (1985) reported that the grain storage pest Plodia interpunctella developed resistance to the spore-crystal complex of B. thuringiensis; after 15 generations of selection with the Indian meal moth, Plodia interpunctella, using a commercial thuringiensis preparation ("Dipel", HD-1 Laboratories, North Chicago, Illinois 60064, USA), a 100-fold decrease in B. thuringiensis sensitivity was reported. Each of the colonies was cultured for several generations on a diet treated with a constant B. thuringiensis dosage which was expected to produce 70-90% larval mortality. Under these high selection presssure conditions, insect resistance to <u>B</u>.

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thuringiensis increased rapidly. More recently, development of resistance against B. thuringiensis is also reported for the almond moth, <u>Cadra cautella</u> (McGaughey and Beeman, 1988). Resistance was stable when selection was discontinued and was inherited as a recessive trait (McGaughey and Beeman, 1988). The mechanism of insect resistance to <u>B. thuringiensis</u> toxins of <u>Plodia interpunctella</u> and <u>Cadra cautella</u> has not been elucidated.

The main cause of <u>B</u>. <u>thuringiensis</u> resistance development in both reported cases involving grain storage was the environmental conditions prevailing during the grain storage. Under the conditions in both cases, the environment was relatively stable, so <u>B</u>. <u>thuringiensis</u> degradation was slow and permitted successive generations of the pest to breed in the continuous presence of the microbial insecticide. The speed at which <u>Plodia</u> developed resistance to <u>B</u>. <u>thuringiensis</u> in one study suggests that it could do so within one single storage season in the bins of treated grain.

Although insect resistance development against B. thuringiensis has mostly been observed in laboratory and pilot scale studies, very recent indications of B. thuringiensis resistance development in xylostella populations in the (cabbage) field have been reported (Kirsch and Schmutterer, 1988). A number of factors have led to a continuous exposure of P. xylostella to B. thuringiensis in a relatively small geographic area. This and the short generation cycle of P. xylostella have seemingly led to an enormous selection pressure resulting in decreased susceptibility and increased resistance thuringiensis.

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A procedure for expressing a B. thuringiensis ICP gene in plants in order to render the plants insectresistant (European patent publication ("EP") 0193259 [which is incorporated herein by reference]; Vaeck et al., 1987; Barton et al., 1987; Fischhoff et al., 1987) provides an entirely new approach to insect control in agriculture which is at the same time environmentally attractive and cost-effective. important determinant for the success of this approach will be whether insects will be able to develop resistance to B. thuringiensis ICPs expressed in transgenic plants (Vaeck et al., 1987; Barton et al., 1987; Fischhoff et al., 1987). In contrast with a foliar application, after which B. thuringiensis ICPs are rapidly degraded, the transgenic plants will exert a continuous selection pressure. It is clear from laboratory selection experiments that a continuous selection pressure has led to adaptation to thuringiensis and its components in several insect species. In this regard, it should be pointed out that the conditions in the laboratory which resulted in the development of insect-resistance to B. thuringiensis are very similar to the situation with transgenic plants which produce B. thuringiensis ICPs and provide a continuous selection pressure on insect populations feeding on the plants. Mathematical models of selection pressure predict that, if engineered insect-resistant plants become a permanent part of their environment, resistance development in insects will emerge rapidly (Gould, 1988). Thus, the chances for the development of insect resistance to B. thuringiensis in transgenic plants may be considerably increased as compared to the field application of B. thuringiensis sprays. Heliothis virescens strain has been reported that is 20 times more resistant to B. thuringiensis HD-1 ICP

produced by transgenic <u>Pseudomonas fluorescens</u> and 6 times more resistant to the pure ICP (Stone et al., 1989). Furthermore, the monetary and human costs of resistance are difficult to assess, but loss of pesticide effectiveness invariably entails increased application frequencies and dosages and, finally, more expensive replacement compounds as new pesticides become more difficult to discover and develop.

Therefore, it would be desirable to develop means for delaying or even preventing the evolution of resistance to <u>B. thuringiensis</u>.

B. thuringiensis strains, active against Lepidoptera (Dulmage et al., 1981), Diptera (Goldberg and Margalit, 1977) and Coleoptera (Krieg et al., 1983), have been described. It has become clear that there is a substantial heterogeneity among ICPs from different strains active against Lepidoptera, as well as among ICPs from strains active against Coleoptera (Hofte and Whiteley, 1989). An overview of the different B. thuringiensis ICP genes, that have been characterized, is given in Table 2 (which follows the Examples herein).

Most of the anti-Lepidopteran B. thuringiensis (e.g., Bt3, Bt2, Bt73, Bt14, Bt15, Bt4, Bt18) ICP genes encode 130 to 140 kDa protoxins which dissolve in the alkaline environment of an insect's midgut and are proteolytically activated into an active toxin of 60-65 kDa. These ICPs are related and can be recognized as of the same family based divergence however homologies. The sequence substantial, and the insecticidal spectrum, among the order Lepidoptera, may be substantially different (Höfte et al., 1988).

The P2 toxin gene and the cry B2 gene are different from the above-mentioned genes in that they

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do not encode high molecular weight protoxins but rather toxins of around 70 kDa (Donovan et al., 1988 and Widner and Whiteley, 1989, respectively).

It has recently become clear that heterogeneity exists also in the anti-Coleopteran toxin gene family. Whereas several previously reported toxin gene sequences from different <u>B. thuringiensis</u> isolates with anti-Coleopteran activity were identical (EP 0149162 and 0202739), the sequences and structure of bt21 and bt22 are substantially divergent (European patent application ("EPA") 89400428.2).

While the insecticidal spectra of B. thuringiensis ICPs are different, the major pathway of their toxic action is believed to be common. All B. thuringiensis ICPs, for which the mechanism of action has been studied in any detail, interact with the midgut epithelium of sensitive species and cause lysis of the epithelial cells (Knowles and Ellar, 1986) due to the fact that the permeability characteristics of the brush border membrane and the osmotic balance over this membrane are perturbed. In the pathway of toxic action of  $\underline{B}$ . thuringiensis ICPs, the binding of the toxin to receptor sites on the brush border membrane of these cells is an important feature (Hofmann et al., 1988b). The toxin binding sites in the midgut can be regarded as an ICP-receptor since toxin is bound in a saturable way and with high affinity (Hofmann et al., 1988a).

Although this outline of the mode of action of <u>B</u>. thuringiensis ICPs is generally accepted, it remains a matter of discussion what the essential determinant(s) are for the differences in their insecticidal spectra. Haider et al. (1986) emphasize the importance of specific proteases in the insect midgut. Hofmann et al. (1988b) indicate that receptor binding is a prerequisite for toxic activity and describe that

<u>Pieris brassicae</u> has two distinct receptor populations for two toxins. Other authors have suggested that differences in the environment of the midgut (e.g., pH of the midgut) might be crucial.

### SUMMARY OF THE INVENTION

In accordance with this invention, a plant is provided having, stably integrated into its genome, at least two <u>B</u>. <u>thuringiensis</u> ICP genes encoding at least two non-competitively binding insecticidal <u>B</u>. <u>thuringiensis</u> ICPs, preferably the active toxins thereof, against a specific target insect, preferably against a Lepidoptera or Coleoptera. Such a plant is characterized by the simultaneous expression of the at least two non-competitively binding <u>B</u>. <u>thuringiensis</u> ICPs.

Also in accordance with this invention, at least two ICP genes, particularly two genes or parts thereof non-competitively binding two coding for Lepidopteran or anti-Coleopteran B. thuringiensis ICPs, are cloned into a plant expression vector. Plant cells transformed with this vector are characterized by the simultaneous expression of the at least two thuringiensis ICP genes. The resulting transformed plant cell can be used to produce a transformed plant in which the plant cells: 1. contain the at least two B. thuringiensis ICP genes or parts thereof encoding at least two non-competitively binding anti-Lepidopteran or anti-Coleopteran B. thuringiensis ICPs as a stable insert into their genome; and 2. express the genes simultaneously, thereby conferring on improved resistance to at least one target species of insect, so as to prevent or delay development of resistance to B. thuringiensis of the at least one target species of insect feeding on the transformed plant.

Further in accordance with this invention, plant expression vectors are provided which allow integration simultaneous expression of at least two thuringiensis ICP genes in a plant cell and which comprise one or more chimeric genes, each containing in same transcriptional unit: a promoter which functions in the plant cell to direct the synthesis of mRNA encoded by one of the ICP genes; one or more different ICP genes, each encoding a non-competitively binding B. thuringiensis ICP; preferably a marker gene; a 3' non-translated DNA sequence which functions in the plant cell for 3' end formation and the addition of polyadenylate nucleotides to the 3'end of the mRNA; and optionally a DNA sequence encoding a proteasepart between any two ICP genes. sensitive protein

# DETAILED DESCRIPTION OF THE INVENTION Definitions

As used herein, "B. thuringiensis ICP" (or "ICP") understood as an intact protein or a should be has insecticidal activity thereof which part and which can produced in nature by be thuringiensis. An ICP can be a protoxin, as well as an active toxin or another insecticidal truncated part of a protoxin which need not be crystalline and which need not be a naturally occurring protein. In chimaeric toxin this regard, an ICP can be a encoded by the combination of two regions of two different ICP genes as disclosed in EP 0228838.

As used herein, "protoxin" should be understood as the primary translation product of a full-length gene encoding an ICP.

As used herein, "toxin", "toxic core" or "active toxin" should all be understood as a part of a protoxin

which can be obtained by protease (e.g., by trypsin) cleavage and has insecticidal activity.

As used herein, "gene" should be understood as a full-length DNA sequence encoding a protein (e.g., such as is found in nature), as well as a truncated fragment thereof encoding at least the active part (i.e., toxin) of the protein encoded by the full-length DNA sequence, preferably encoding just the active part of the protein encoded by the full-length DNA sequence. A gene can be naturally occurring or synthetic.

As used herein, "truncated  $\underline{B}$ . thuringiensis gene" should be understood as a fragment of a full-length  $\underline{B}$ . thuringiensis gene which still encodes at least the toxic part of the  $\underline{B}$ . thuringiensis ICP, preferentially the toxin.

As used herein, "marker gene" should be understood as a gene encoding a selectable marker (e.g., encoding antibiotic resistance) or a screenable marker (e.g., encoding a gene product which allows the quantitative analysis of transgenic plants).

Two ICPs are said to be "competitively binding ICPs" for a target insect species when one ICP competes for all ICP receptors of the other ICP, which receptors are present in the brush border membrane of the midgut of the target insect species.

Two ICPs are said to be "non-competitively binding ICPs" when, for at least one target insect species, the first ICP has at least one receptor for which the second ICP does not compete and the second ICP has at least one receptor for which the first ICP does not compete, which receptors are present in the brush border membrane of the midgut of the target insect species.

A "receptor" should be understood as a molecule, to which a ligand (here a B. thuringiensis ICP,

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preferably a toxin) can bind with high affinity (typically a dissociation constant (Kd) between 10.11 and 10-6M) and saturability. A determination of whether two ICPs are competitively or non-competitively binding ICPs can be made by determining whether: 1. a first ICP competes for all of the receptors of a second ICP when all the binding sites of the second ICP with an affinity in the range of about 10<sup>-11</sup> to 10<sup>-6</sup>M can be saturated with the first ICP in concentrations of the first ICP of about 10<sup>-5</sup>M or less (e.g., down to about 10-11M); and 2. the second ICP competes for the all of the receptors of the first ICP when all the binding sites of the first ICP with an affinity in the range of about 10-11 to 10-6M can be saturated with the second ICP in concentrations of the second ICP of about 10<sup>-5</sup>M or less.

### General Procedures

This section describes in broad terms general procedures for the evaluation and exploitation of at least two <u>B</u>. <u>thuringiensis</u> ICP genes for prevention of the development, in a target insect, of a resistance to the <u>B</u>. <u>thuringiensis</u> ICPs expressed in transgenic plants of this invention. A non-exhaustive list of consecutive steps in the general procedure follows, after which are described particular Examples that are based on this methodology and that illustrate this invention.

In accordance with this invention, specific <u>B</u>. thuringiensis ICPs can be isolated in a conventional manner from the respective strains such as are listed in Table 2 (which follows the Examples). The ICPs can be used to prepare monoclonal or polyclonal antibodies specific for these ICPs in a conventional manner (Höfte et al., 1988).

The ICP genes can each be isolated from their conventional strains in а respective Preferably, the ICP genes are each identified by: digesting total DNA from their respective strains with suitable restriction enzyme(s); size fractionating the DNA fragments, so produced, into DNA fractions of 5 to 10 Kb; ligating such fractions to suitable cloning vectors (e.g., pEcoR251, deposited at the Deutsche Sammlung von Mikroorganismen und Zellculturen ("DSM"), Braunschweig, Federal Republic of Germany, 4711 July 13, 1988); accession number no. on transforming E.coli with the cloning vectors; screening the clones with a suitable DNA probe. The DNA probe can be constructed from a highly conserved region which is commonly present in different B. thuringiensis genes which encode crystal protoxins against Coleoptera or Lepidoptera, such as on the basis of an N-terminal amino acid sequence determined by gas-phase sequencing of the purified proteins (EPA 88402115.5).

Alternatively, the desired fragments, prepared from total DNA of the respective strains, can be ligated in suitable expression vectors (e.g., a pUC vector (Yanisch-Perron et al., 1985) with the insert under the control of the lac promoter) and transformed in <u>E. coli</u>, and the clones can then be screened by conventional colony immunoprobing methods (French et al., 1986) for expression of the toxins with monoclonal or polyclonal antibodies raised against the toxins produced by the strains.

The isolated <u>B. thuringiensis</u> ICP genes can then be sequenced in a conventional manner using well-known procedures (e.g., Maxam and Gilbert, 1980).

At present, several ICP genes have been cloned from different subspecies of  $\underline{B}$ . thuringiensis (Table 2). The nucleotide sequences from several of these  $\underline{B}$ .

thuringiensis ICP genes have been reported. Whereas several sequences are identical or nearly identical and represent the same gene or slight variants of the same gene, several sequences display substantial heterogeneity and show the existence of different B. thuringiensis ICP gene classes. Several lines of evidence suggest that all these genes specify a family of related insecticidal proteins. Analysis of the distribution of B. thuringiensis ICPs in different B. thuringiensis strains by determining the protein composition of their crystals, by immunodetection using polyclonal antisera or monoclonals against purified crystals, or by using gene-specific probes, shows that subspecies of B. thuringiensis might contain up to three related B. thuringiensis ICP genes belonging to different classes (Kronstad et al., 1983).

To express the isolated and characterized gene in a heterologous host for purification and characterization of the recombinant protein, the preferred organism is Escherichia coli. A number of expression vectors for enhanced expression heterologous genes in E. coli have been described (e.g., Remaut et al., 1981). Usually the gene is cloned under control of a strong regulatable promoter, such as the lambda pL or pR promoters (e.g., Botterman and Zabeau, 1987), the lac promoter (e.g., Fuller, 1982) or the tac promoter (e.g., De Boer et al., 1983), and provided with suitable translation initiation sites (e.g., Stanssens et al, 1985 and 1987). Gene cassettes of the B. thuringiens 3 ICP genes can be generated by site-directed mutagenesis, for example according to the procedure described by Stanssens et al. (1985 and This allows cassettes to be made comprising, for example, a truncated ICP gene fragment encoding the toxic core (i.e., toxin) of an ICP or a hybrid gene

encoding the toxic core and a selectable marker according to the procedures described in EPA 88402241.9.

The cells of an E. coli culture, which has been induced to produce a recombinant ICP, are harvested. The method used to induce the cells to produce the recombinant ICP depends on the choice of the promoter. For example, the lac promoter (Fuller, 1982) is induced by isopropyl-B-D-thiogalacto-pyranoside ("IPTG"); the pL promoter is induced by temperature shock (Bernard et al., 1979). The recombinant ICP is usually deposited in the cells as insoluble inclusions (Hsuing and Becker, 1988). The cells are lysed to liberate the inclusions. The bulk of E. coli proteins is removed in subsequent washing steps. A semi-purified protoxin pellet is obtained, from which the protoxin can be dissolved in alkaline buffer (e.g., Na2CO3, pH 10). The procedure for the ICP Bt2, which is also applicable to other recombinant toxins, has been described by Höfte et al., 1986.

In accordance with this invention, the binding of various ICPs to ICP receptors on the brush border membrane of the columnar midgut epithelial cells of various insect species has been investigated. The brush border membrane is the primary target of each ICP, and membrane vesicles, preferentially derived from the brush border membrane, can be obtained according to Wolfersberger et al., 1987.

The binding to ICP receptors of one or more ICPs (e.g., ICP A, ICP B, etc.) can be characterized by the following steps (Hofmann et al, 1988b):

- 1. ICP A is labelled with a suitable marker (usually a radioisotope such as  $^{125}I$ ).
- 2. Brush border membranes are incubated with a small amount (preferably less than 10<sup>-10</sup> M) of labelled

ICP A together with different concentrations of non-labelled ICP A (preferably from less than  $10^{-11}$  to  $10^{-5}$  M).

- 3. For all concentrations tested the amount of labelled ICP A bound to the brush border membranes is measured.
- 4. Mathematical analysis of these data allows one to calculate various characteristics of the ICP receptor such as the magnitude of the population of binding sites (Scatchard, 1949).
- 5. Competition by other toxins (e.g. ICP B) is preferably studied by incubating the same amount of labelled ICP A with brush border membranes in combination with different amounts of ICP B (preferentially from 10<sup>-11</sup> to 10<sup>-6</sup> M; and subsequently, steps 3 and 4 are repeated.

By this procedure, it has been found, for example, that Bt3 toxin, Bt2 toxin and Bt73 toxin are competitively binding anti-Lepidopteran ICPs for Manduca sexta and Heliothis virescens (See example 6 which follows). Various other combinations of toxins have been found to be non-competitively binding anti-Lepidopteran or anti-Coleopteran toxins (example 6).

Although the concept of competitivity versus noncompetitivity of ICP binding does not have practical importance by itself, the observation of the non-competitivity of two  $\underline{B}$ . thuringiensis ICPs, active against the same target insect, can be put to very significant practical use. This is because combination of two non-competitively binding thuringiensis ICPs can be used to prevent development, by a target insect, of resistance against such B. thuringienis ICPs.

A selection experiment with  $\underline{M}$ .  $\underline{sexta}$ , using Bt2 toxin, Bt18 toxin, and a mixture of Bt2 and Bt18

toxins, has shown that Bt2 and Bt18 are two noncompetitively binding anti-Lepidopteran toxins. After very pronounced generations of selection, a reduction in ICP sensitivity was observed in the selection experiments with Bt2 or Bt18 alone (>100 times). The reduction in sensitivity in the selection experiment with a Bt2-Bt18 mixture was only marginal (3 times). This demonstrates the unexpected practical simultaneous use of two advantage of a competitively binding ICPs in a situation which models the high selection pressure which will exist with the use of transgenic plants transformed with ICP genes. In this regard, the two resistant strains showed a specific loss in receptor sites for either the Bt2 or Bt18 toxin. In each case, receptor sites for the toxin, which was not used for selection, were not affected or their concentration even increased. Thus, the Bt2 selected strain retained its Bt18 receptors, and the Bt18 selected strain developed an increased number of Bt2 receptors. Indeed, the Bt18 selected strain showed increased sensitivity for Bt2 along with its increased Bt2 receptor concentration. No significant changes in receptor sites were found in the strain selected against the combined toxins. These findings are described in detail in Example 7 which follows.

A similar mechanism of resistance to Bt has been observed with respect to a strain of diamondback moth, Plutella xylostella. This strain had developed resistance in the field to Dipel which is a commercial formulation of the Bt HD-1 strain. Crystals of Dipel comprise a mixture of several BtICPs, similar to the Bt2, Bt3 and Bt73 proteins which are competitively-binding ICPs. As shown by both insect bioassays and competitive binding studies using Bt2 and Bt15, the Dipel-resistant diamondback moth strain is resistant to

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Bt2 protoxin and toxin but maintains full sensitivity to Bt15 protoxin and toxin. This finding is relevant to other combinations of non-competitively binding anti-Lepidopteran or Coleopteran ICPs which are expected to have the same beneficial effect against their common target insects.

Hence, a combination of non-competitively binding ICPs, when directly expressed in a transgenic plant, offers the substantial advantage of reducing the chances of development of insect resistance against the ICPs expressed in the plant. There may be additional benefits because the combined spectrum of two toxins may be broader than the spectrum of a single ICP expressed in a plant (See Examples 8, 9 and 10 which follow).

If, among two competitively binding ICPs, one has a larger binding site population than the other against a given target insect, it will be most advantageous to use the one with the larger population of binding sites to control the target pest in combination with the most suitable non-competitively binding B. thuringiensis ICP. For example, as seen from Example 6, it is preferred to use Bt73 against Heliothis virescens, rather than Bt2 or Bt3, and it is preferred to use Bt3 against Manduca sexta rather than Bt2 or Bt73. The selected gene can then be combined with the best suitable non-competitively binding ICP.

Previously, plant transformations involved the introduction of a marker gene together with a single ICP gene, within the same plasmid, in the plant genome (e.g., Vaeck et al., 1987; Fischoff et al., 1987). Such chimeric ICP genes usually comprised either all or part of an ICP gene, preferably a truncated ICP gene fragment encoding the toxic core, fused to a selectable marker gene, such as the neo gene coding for neomycin

phosphotransferase. The chimeric ICP gene was placed between the T-DNA border repeats for Agrobacterium Tiplasmid mediated transformation (EP 0193259).

This invention involves the combined expression of two or even more B. thuringiensis ICP genes The insecticidally effective transgenic plants. thuringiensis ICP genes, encoding two non-competitively binding ICPs for a target insect species, preferably encoding the respective truncated ICP genes, inserted in a plant cell genome, preferably in its nuclear genome, so that the inserted genes downstream of, and under the control of, a promoter which can direct the expression of the genes in the This is preferably accomplished plant cell. inserting, in the plant cell genome, one or more chimaeric genes, each containing in the transcriptional unit: at least one ICP gene; preferably a marker gene; and optionally a DNA sequence encoding a (e.g., trypsin) - sensitive or protein part intercalated in frame between any two ICP genes in the chimaeric gene. Each chimaeric gene also contains at least one promoter which can direct expression of its ICP gene in the plant cell.

The selection of suitable promoters for the chimaeric genes of this invention is not critical. Preferred promoters for such chimaeric genes include: the strong constitutive 35S promoter obtained from the cauliflower mosaic virus, isolates CM 1841 (Gardner et al., 1981), CabbB-S (Franck et al., 1980) and CabbB-JI (Hull and Howell, 1987); the promoter of the nopaline synthetase gene ("PNOS") of the Ti-plasmid (Herrera-Estrella, 1983); the promoter of the octopine synthase gene ("POCS" [De Greve et al., 1982]); and the wound-inducible TR1' promoter and the TR2' promoter which drive the expression of the 1' and 2' genes,

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respectively, of the T-DNA (Velten et al., 1984). Alternatively, a promoter can be utilized which is specific for one or more tissues or organs of the plant, whereby the inserted genes are expressed only in cells of the specific tissue(s) or organ(s). Examples of such promoters are a stem-specific promoter such as the AdoMet-synthetase promoter (Peleman et al., 1989), a tuber-specific promoter (Rocha-Sosa et al., 1989), and a seed-specific promoter such as the 2S promoter (Krebbers et al., 1988). The ICP genes could also be selectively expressed in the leaves of a plant (e.g., potato) by placing the genes under the control of a light-inducible promoter such as the promoter of the ribulose-1,5-bisphosphate carboxylase small gene of the plant itself or of another plant such as pea as disclosed in EP 0193259. Another alternative is to use a promoter whose expression is inducible (e.g., by temperature or chemical factors).

A 3' non-translated DNA sequence, which functions in plant cells for 3' end formation and the polyadenylation of the 3' end of the mRNA sequence encoded by the at least one ICP gene in the plant cell, also forms part of each such chimeric gene. The selection of a suitable 3' non-translated DNA sequence is not critical. Examples are the 3' untranslated end of the octopine synthase gene, the nopaline synthase gene or the T-DNA gene 7 (Velten and Schell, 1985).

The selection of marker genes for the chimaeric genes of this invention also is not critical, and any conventional DNA sequence can be used which encodes a protein or polypeptide which renders plant cells, expressing the DNA sequence, readily distinguishable from plant cells not expressing the DNA sequence (EP 0344029). The marker gene can be under the control of its own promoter and have its own 3' non-translated DNA

sequence as disclosed above, provided the marker gene is in the same genetic locus as the ICP gene(s) which it identifies. The marker gene can be, for example: a herbicide resistance gene such as the sfr or sfrv genes (EPA 87400141); a gene encoding a modified target enzyme for a herbicide having a lower affinity for the (non-modified) than the natural herbicide enzyme, such as a modified 5-EPSP as a target for glyphosate (U.S. patent 4,535,060; EP 0218571) or a modified glutamine synthetase as a target for a glutamine synthetase inhibitor (EP 0240972); or an antibiotic resistance gene, such as a neo gene (PCT publication W0 84/02913; EP 0193259).

Using A. tumefaciens Ti vector-mediated plant transformation methodology, all chimeric genes of this invention can be inserted into plant cell genomes after the chimaeric genes have been placed between the T-DNA border repeats of suitable disarmed Ti-plasmid vectors (Deblaere et al., 1988). This transformation can be carried out in a conventional manner, for example as described in EP 0116718, PCT publication WO 84/02913 and EPA 87400544.0. The chimeric genes can also be in non-specific plasmid vectors which can be used for direct gene transfer (e.g., as described by Pazkowski et al., 1984; De La Pena et al., 1986). Different conventional procedures can be followed to obtain a combined expression of two B.thuringiensis ICP genes in transgenic plants as summarized below.

- I Chimeric gene constructs whereby two or more ICP genes and a marker gene are transferred to the plant genome as a single piece of DNA and lead to the insertion in a single locus in the genome
- <u>Ia</u> <u>The genes can be engineered in different transcriptional units each under control of a distinct promoter</u>

To express two or more ICP genes and a marker gene as separate transcriptional units, several promoter fragments directing expression in plant cells can be used as described above. All combinations of the promoters mentioned above in the chimaeric constructs for one ICP gene are possible. Examples of such individual chimeric constructs are described for the bt2 gene in EP 0193259, for the bt13 gene in EPA 88402115.5 and for the bt18 gene in EPA 88402241.9. The ICP gene in each chimeric gene of this invention can be the intact ICP gene or preferably an insecticidallyeffective part of the intact ICP gene, especially a truncated gene fragment encoding the toxic core of the ICP. The individual chimeric genes are cloned in the same plasmid vector according to standard procedures (e.g., EP 0193259).

Ib Two genes (e.g., either an ICP and a marker gene or two ICP genes) or more can be combined in the same transcriptional unit

To express two or more ICP genes in the same transcriptional unit, the following cases can be distinguished:

In a first case, hybrid genes in which the coding region of one gene is in frame fused with the coding region of another gene can be placed under the control of a single promoter. Fusions can be made between either an ICP and a marker gene or between two ICP genes. An example of an ICP gene-marker gene fusion has been described in EP 0193259 (i.e., a hybrid truncated <a href="https://doi.org/bt/>bt2-neo">bt2-neo</a> gene encoding a Bt2 toxin-NPTII fusion protein).

Another possibility is the fusion of two ICP genes. Between each gene encoding an ICP which still is insecticidally active (i.e., a toxic part of the protoxin), a gene fragment encoding a protease (e.g.,

trypsin) - sensitive protein part should be included, such as a gene fragment encoding a part of the N-terminal or C-terminal amino acid sequence of one of the ICPs which is removed or cleaved upon activation by the midgut enzymes of the target insect species.

In a second case, the coding regions of the two respective ICP genes can be combined in dicistronic units placed under the control of a promoter. The coding regions of the two ICP genes are placed after each other with an intergenic sequence of defined length. A single messenger RNA molecule is generated, leading to the translation into two separate gene products. Based on a modified scanning model (Kozak, 1987), the concept of reinitiation of translation has been accepted provided that a termination codon in frame with the upstream ATG precedes the downstream ATG. Experimental data also demonstrated that the plant translational machinery is able to synthesize several polypeptides from a polycistronic mRNA (Angenon et al., 1989).

# II Chimeric constructs with one or more ICP genes that are transferred to the genome of a plant already transformed with a one or more ICP genes

Several genes can be introduced into a plant cell during sequential transformation steps (retransformation), provided that an alternative system to select transformants is available for the second round of transformation. This retransformation leads to the combined expression of ICP genes which are introduced at multiple loci in the genome. Preferably, different selectable marker genes are used in the two consecutive transformation steps. The first marker is used for selection of transformed cells in the first transformation, while the second marker is used for selection of transformants in the second round of

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transformation. Sequential transformation steps using kanamycin and hygromycin have been described, for example by Sandler et al. (1988) and Delauney et al. (1988).

III Chimeric constructs with one or more ICP genes, that are separately transferred to the nuclear genome of separate plants in independent transformation events and are subsequently combined in a single plant genome through crosses.

The first plant should be a plant transformed with a first ICP gene or an F1 plant derived herefrom through selfing (preferably an F1 plant which is homozygous for the ICP gene). The second plant should be a plant transformed with a second ICP gene or an F1 plant derived herefrom through selfing (preferably an F1 plant which is homozygous for the second ICP gene). Selection methods can be applied to the plants obtained from this cross in order to select those plants having the two ICP genes present in their genome (e.g., Southern blotting) and expressing the two ICPs (e.g., separate ELISA detection of the immunologically different ICPs). This is a useful strategy to produce hybrid varieties from two parental transformed with a different ICP gene, as well as to produce inbred lines containing two different ICP genes through crossing of two independent transformants (or their F1 selfed offspring) from the same inbred line. IV Chimeric constructs with one or more ICP genes separately transferred to the genome of a single plant in the same transformation experiment leading to the insertion of the respective chimeric genes at multiple loci.

Cotransformation involves the simultaneous transformation of a plant with two different expression vectors, one containing a first ICP gene, the second

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containing a second ICP gene. Along with each ICP gene, a different marker gene can be used, selection can be made with the two simultaneously. Alternatively, a single marker can be used, and a sufficiently large number of selected plants can be screened in order to find those plants having the two ICP genes (e.g., by Southern blotting) and expressing the two proteins (e.g., by means of ELISA). Cotransformation with more than one T-DNA can be accomplished by using simultaneously two different strains of Agrobacterium, each with a different Tiplasmid (Depicker et al., 1985) or with one strain of Agrobacterium containing two T-DNAs on separate plasmids (de Framond et al., 1986). Direct gene transfer, using a mixture of two plasmids, can also be employed to cotransform plant cells with a selectable and a non-selectable gene (Schocher et al., 1986).

The transgenic plant obtained can be used in further plant breeding schemes. The transformed plant can be selfed to obtain a plant which is homozygous for the inserted genes. If the plant is an inbred line, this homozygous plant can be used to produce seeds directly or as a parental line for a hybrid variety. The gene can also be crossed into open pollinated populations or other inbred lines of the same plant using conventional plant breeding approaches.

Of course other plant transformation methods can be used and are within the scope of the invention as long as they result is a plant which expresses two or more non-competitively binding ICPs. In this regard, this invention is not limited to the use of Agrobacterium Ti-plasmids for transforming plant cells with genes encoding non-competitively binding ICPs. Other known methods for plant cell transformations, such as electroporation or by the use of a vector

system based on plant viruses or pollen, can be used for transforming monocotyledonous and dicotyledonous plants in order to obtain plants which express two non-competitively binding ICPs. Furthermore. DNA sequences encoding two non-competitively binding ICPs other than those disclosed herein can be used for transforming plants. Also, each of the ICP genes, described herein, can be encoded by equivalent DNA sequences, taking into consideration the degeneracy of the genetic code. Also, equivalent ICPs with only a few amino acids changed, such as would be obtained through mutations in the ICP gene, can also be used, provided they encode a protein with essentially the same characteristics (e.g., insecticidal activity and receptor binding).

The following Examples illustrate the invention. Those skilled in the art will, however, recognize that other combinations of two or more non-competitively binding B. thuringiensis ICP genes can be used to transform plants in accordance with this invention in order to prevent the development, in a target insect, of resistance to B. thuringiensis ICPs expressed in the transformed plants. Unless otherwise indicated, all procedures for making and manipulating DNA were carried out by the standardized procedures described in Maniatis et al, Molecular Cloning - A Laboratory Manual, Cold Spring Harbor Laboratory (1982).

### EXAMPLE 1: Collection of genes

The collection of anti-Lepidopteran and anti-Coleopteran Bt genes encoding ICPs, which are the subject of the Examples, is described in Table 2 (following the Examples). References for the respective genes are indicated in Table 2. The origin, the isolation and characterization of the Bt genes, which have not been published, are described below. Bt

strains, such as strains HD-1, HD-68, HD-110, and HD-73, are publicly available from the Agricultural Research Culture Collection, Northern Regional Research Laboratory, U.S. Dept. of Agriculture, Peoria, Illinois 61604, U.S.A.

bt3

gene: From <u>B</u>. <u>thuringiensis</u> var. kurstaki HD-1, the ICP was cloned. Characterization of this gene revealed an open reading frame of 3528 bp which encodes a protoxin of 133 kDa. This gene was identical to the one described by Schnepf et al. (1985).

**bt73** 

gene: From <u>B. thuringiensis</u> var HD-73. The ICP gene was cloned as described by Adang et al. (1985).

bt4

gene: A genomic library was prepared from total DNA of strain B. thuringiensis aizawai HD-68. Using the 1.1 kb internal HindIII fragment of the bt2 gene as a probe, a gene designated bt4 was isolated. Characterization of this gene revealed an open reading frame of 3495 bp which encodes a protoxin of 132 kDa and a trypsin activated toxin fragment of 60 kDa. This (insect controlling protein) gene differs from previously identified genes and was also found in several other strains of subspecies aizawai and entomocidus including HD-110. Fig. 13 shows the nucleotide sequence and deduced amino acid sequence of the open reading frame ("ORF") of the bt4 gene extending from nucleotide 264 to nucleotide 3761.

bt14 and bt15

genes: A genomic library was prepared from total DNA of strain B. thuringiensis var. entomocidus HD-110 by partial Sau3A digest of the total DNA and cloning in the vector pEcoR251 (deposited at DSM under

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accession number 4711). Using monoclonal antibodies (Höfte et al., 1988), at least three structurally distinct ICPs were identified in crystals of B. thuringiensis entomocidus HD-110. These monoclonal antibodies were used to clone the three different ICP genes from this B. thuringiensis strain. One of these genes is the bt4 gene as described above.

The second gene was called "bt15". Fig. 14 shows the nucleotide sequence and deduced amino acid sequence of the ORF of the bt15 gene, isolated from HD-110, extending from nucleotide 234 to nucleotide 3803. The Shine and Dalgarno sequence, preceding the initiation codon is underlined. This gene has an open reading frame of 3567 bp which encodes a protoxin of 135 kDa and a 63 kDa toxin fragment. A similar gene has been described by Honee et al. 1988, isolated from B. thuringiensis entomocidus 60.5. The bt15 gene differs from the published sequence at three positions: an Ala codon (GCA) is present instead of an Arg codon (CGA) at position 925 and a consecution of a Thr-His codon (ACGCAT) is present instead of a Thr-Asp codon (ACCGAT) at position 1400. (The numbers of the positions are according to Honnee et al., 1988). Another similar gene has been described in EP 0295156, isolated from B. thuringiensis aizawai 7-29 and entomocidus 6-01. The bt15 gene is different from this published nucleotide sequence at three different places : 1) a Glu codon (GAA) instead of an Ala codon (GCA) at position 700; 2) the sequence TGG, CCA, GCG, CCA instead of TGC, CAG, CGC, CAC, CAT at position 1456 and 3) an Arg codon (CGT) instead of an Ala codon (GCG) at

position 2654. (The numbers of the positions are according to EP 0295156).

The third gene isolated was called "bt14". It has an open reading frame of 3621 bp which encodes a 137 kDa protoxin and a 66 kDa activated toxin fragment. A similar gene has been cloned from B.thuringiensis HD-2 (Brizzard and 1988). The bt14 gene differs from the published nucleotide sequence by two nucleotide substitutions: a T instead of a C at position 126, and a C instead of a T at position 448 (the numbers of the positions are according to Brizzard and Whiteley, 1988). In the first case, the Ile codon (ATT or ATC) is conserved whereas in the second case the Tyr codon (TAT) is converted to a His codon (CAC).

bt2

gene: The <u>bt2</u> gene was cloned as described in EP 0193259.

bt18

gene: Cloning of the <u>bt18</u> gene was performed as described in EPA 88402241.9.

bt13

gene: The <u>bt13</u> gene was cloned as described in EPA 88402115.5.

bt21 and bt22

genes: These genes, encoding Coleopteran-active ICPs, were cloned as described in EPA 89400428.2.

# EXAMPLE 2 : Construction of gene cassettes and expression of Bt genes in E.coli

1) bt2, bt18: the construction of bt2 and bt18 gene cassettes has been previously described in EPA 86300291.1 and 88402241.9, respectively. Basically, they comprise a truncated gene encoding the toxic core and a hybrid gene comprising the

truncated gene fused in frame to the N-terminus of the  $\underline{\text{neo}}$  gene. The gene cassettes are used to transform  $\underline{\text{E}}$ .  $\underline{\text{coli}}$  to express the Bt2 and Bt18 ICP toxins.

2) bt14, bt15: as described in EPA 88402241.9, gene cassettes for the <u>bt14</u> and <u>bt15</u> genes were constructed in order to express the genes in <u>E.coli</u> and in plants.

First, a NcoI site was introduced at the N-terminus of the genes by site-directed mutagenesis.

In the case of the <u>bt15</u> gene, the conversion of the TT nucleotides, immediately in front of the ATG codon, into CC yielded a NcoI site overlapping with the ATG initiation codon. This site was introduced using the pMa/c vectors for site-directed mutagenesis (Stanssens et al., 1987) and a 28-mer oligonucleotide with the following sequence:

5'-CGGAGGTATTCCATGGAGGAAAATAATC-3'.

This yielded the plasmid pVE29 carrying the N-terminal fragment of the <u>bt15</u> gene with a NcoI site at the ATG initiation codon.

According to Brizzard and Whiteley (1988), the initiation codon of the <u>bt14</u> gene is a TTG codon. Thus, a NcoI site was created in a like manner at this codon for site directed mutagenesis using a 34-mer oligonucleotide with the following sequence: 5'-CCTATTTGAAGCCATGGTAACTCCTCCTTTTATG-3'.

In this case the sequence of the intitiation codon was converted from ATATTGA to ACCATGG. This yielded the plasmid pHW44 carrying the N-terminal fragment of the <a href="https://doi.org/10.1007/bb14">bt14</a> gene with a NcoI site at the initiation codon.

In a second step, the genes were reconstructed by ligating the N-terminal gene fragments with a suitable C-terminal gene fragment, yielding a <u>bt15</u> gene and <u>bt14</u> gene with a NcoI site at the ATG initiation codon.

To express the <u>bt14</u> and <u>bt15</u> genes encoding the protoxin in <u>E. coli</u>, the following constructs were made: pOH50 containing the <u>bt15</u> gene under the control of the lac promoter; and pHW67 containing the <u>bt14</u> gene under the control of the tac promoter. Induction of a culture of the <u>E. coli</u> strain WK6 carrying the respective plasmids with IPTG yielded an overproduced protein (Fuller, 1982).

The active toxic fragments of the Bt15 and Bt14 protoxins comprise 63 and 60 kDa trypsin digest products respectively. Instead of expressing the whole <a href="https://doi.or/bt14">bt15</a> or <a href="https://doi.or/bt14">bt14</a> gene, it is also possible to express a toxin-encoding gene fragment or derivative thereof in plants. To this end, truncated <a href="https://doi.or/bt14">bt14</a> and <a href="https://doi.or/bt15">bt15</a> gene fragments were constructed. In order to be able to select transgenic plants producing the ICP gene products, hybrid genes of the truncated gene fragments were also made with the <a href="mailto:neo-gene">neo-gene</a> encoding a selectable marker as described in EP 0193259.

By comparison of the nucleotide sequence of the bt4, bt14 and bt15 genes, respectively, with the bt2 and bt18 genes, respectively, the BclI site could be identified as a suitable site localized downstream of the coding sequence encoding the toxin gene fragment. To construct a truncated gene fragment and a hybrid gene of the truncated gene fragment with the neo gene, the filled BclI site was ligated to the filled EcoRI site of pLKM91 (Höfte et al., 1986) and the filled HindIII site of pLK94 respectively (Botterman Zabeau, 1987). pLKM91 carries a 5' truncated neo gene fragment which codes for an enzymatically active Cterminal gene fragment of the neo gene, and pLK94 contains translation stop codons in three reading frames. This yielded the following plasmids which are then used to transform E. coli to express the ICP genes: pHW71 carrying a truncated <u>bt14-neo</u> hybrid gene; pHW72 carrying a truncated <u>bt14</u> gene; pVE34 carrying a truncated <u>bt15-neo</u> hybrid gene; and pVE35 carrying a truncated bt15 gene.

In a similar way as described for the  $\underline{bt14}$  and  $\underline{bt15}$  genes, gene cassettes are constructed for the  $\underline{bt3}$  and  $\underline{bt4}$  genes which are then expressed in  $\underline{E.coli}$ .

### EXAMPLE 3: Purification of recombinant ICPs .

The ICPs expressed in E. coli in Example 2 are purified by the method (described for recombinant Bt2 protoxin) by Höfte et al. (1986).

### EXAMPLE 4: Purification of toxins

Solubilized protoxins of Bt2, Bt3, Bt73, Bt4, Bt14, Bt15, Bt18, Bt13, Bt21 and Bt22 (in  $Na_2CO_3$  50mM, DTT 10 mM pH=10) are dialyzed against 0.5 % ( $NH_4$ )<sub>2</sub>CO<sub>3</sub> at pH 8 and treated with trypsin (trypsin/protoxin=1/20 w/w) for 2h at 37°C. The activated toxin is chromatographically purified (Mono-Q column on FPLC) as described by Hofmann et al.(1988b).

### EXAMPLE 5: Determination of the insecticidal spectrum

The ICP protoxins and toxins of Examples 3 and 4 are evaluated for their insecticidal activity. Each protoxin is dissolved in alkaline buffer containing a reducing agent (Na<sub>2</sub>CO<sub>3</sub> 50 mM, DTT 10 mM pH=10), and each toxin is used as soluble protein directly from FPLC. Protein concentrations are determined. Subsequently, dilutions of the resulting protoxin or toxin solution are prepared in PBS buffer pH=7.4 containing 0.15 M NaCl and 0.1 % bovine serum albumin ("BSA").

The artificial medium for insect culture, described by Bell and Joachim (1976) for Manduca sexta, is poured in appropriate receptacles and allowed to solidify. Subsequently a quantity of the (pro)toxin dilutions is applied on this medium, and the water is

allowed to evaporate under a laminar flow. This results in a medium with a certain quantity (in the range of 0.1 to 10000 ng/cm2) of toxin coated on its surface. For example, for the Bt2 toxin, typical dilutions for a toxicity test on Manduca sexta are 1, 5, 25, 125 and 625 ng/cm2. First instar larvae of Manduca sexta are then applied on the coated medium, and growth and mortality are assessed after 6 days. Mortality increases with dosage. Dose response data is analysed in probit analysis (Finney, 1962), and the data are best summarized by an LD50 value which is the amount of toxin which kills 50 % of the insects. The LD50 for Bt2 toxin against Manduca sexta is around 20 ng/cm2.

Similar assays are carried out for other insect species using a suitable diet or by applying the ICPs on leaves for insects, for which no artificial diet is used.

### EXAMPLE 6: Binding studies

#### Toxins

All protoxins and their toxic fragments were purified according to the methods described for the Bt2 protoxin and toxin in Höfte et al. (1986) and EP 0193259. The activated and purified toxins are further referred to as the Bt2, Bt3, Bt73, Bt4, Bt14, Bt15, Bt18, Bt13, Bt21 and Bt22 toxins.

By way of example for the Bt73 toxin, it has been shown that B. thuringiensis var. kurstaki HD73 produces a protein of 133 kDa encoded by a 6.6 kb type gene. A culture of this strain was grown as described by Mahillon and Delcour (1984). The autolysed culture was spun down (20 minutes at 4500 rpm in a HB4 rotor) and washed with a buffer containing 20 mM Tris, 100 mM NaCl and 0.05 % Triton X-100, pH 8. The final pellet was resuspended in this buffer (4 ml buffer for 100 ml culture). This solution was then layered onto a linear

Urograffin gradient (60-70%) which was centrifuged in a SW 28 rotor for 90 minutes at 18000 rpm. Crystals were collected and stored at -20° C until further use. Activation was performed according to Höfte et al. (1986). The purified toxin is further referred to as the Bt73 toxin.

### Iodination of ICPs

Iodination of Bt2, Bt3, and Bt73 toxins was performed using the Chloramin-T method (Hunter and Greenwood, 1962). 1 mCi  $^{125}$ I-NaI and 20 to 37.5 ug Chloramin-T in NaCl/P; were added to 50 ug of purified toxin. After gentle shaking for 60 seconds, the reaction was stopped by adding 53 ug of potassium metabisulfite in  $H_2O$ . The whole mixture was loaded on a PD 10 Sephadex G-25M gelfiltration column to remove free iodine. A subsequent run on a Biogel P-60 column was carried out in order to increase the purity.

Alternatively, toxins were labeled using Iodogen method. Iodogen (Pierce) was dissolved in chloroform at 0.1 mg/ml. 100 ul of this solution was pipetted into a disposable glass vessel and dried under The vessel was rinsed with a stream of nitrogen gas. Tris buffer (20 mM Tris, pH 8.65 with 0.15 M NaCl). ug of toxin (in Tris buffer) was incubated with 1 mCi of 125I-NaI in the tube for 10 minutes. The reaction was then stopped by the addition of 1 M NaI ( one fourth of the sample volume). The sample was immediately loaded onto a PD10 Sephadex G-25M column and later on a Biogel P-60 column to remove free iodine and possible degradation products.

Other toxins were iodinated using one of the above mentioned procedures.

### Determination of specific activity of iodinated toxin

Specific activity of iodinated Bt2, Bt3, and Bt73 toxin samples was determined using a "sandwich" ELISA

technique according to Voller, Bidwell and Barlett (1976). Primary antibody was a polyclonal antiserum raised against Bt2 toxin, and the secondary antibody was a monoclonal antibody 4D6.

conjugate used was The alkaline phosphatase coupled to anti-mouse IgG. The reaction intensity of a standard dilution series of unlabeled toxin and dilutions of the iodinated toxin sample (in NaCl/P, -BSA) measured. was Linear regression calculations yielded the protein content of radioactive toxin sample. The samples with the highest specific activities were used in the binding assays. Specific activities were 59400, 33000 and 19800 Ci/mole (on reference date) for Bt73 toxin (labeled according to Iodogen procedure), Bt2 toxin (Chloramin-T method) and Bt3 toxin (Iodogen method) respectively.

Specific activities of other toxins were determined using a similar approach. Specific monoclonal and polyclonal antibodies for each of these toxins were raised and applied in ELISA.

### Preparation of brush border membrane vesicles

Brush border membrane vesicles ("BBMV") Manduca sexta, Heliothis virescens, Plutella xylostella, Phthorimaea operculella, Spodoptera exigua, Spodoptera littoralis, Plodia interpunctella, Mamestra brassicae, Pieris brassicae and Leptinotarsa decemlineata were prepared according to the method of Wolfersberger et al. (1987). This is a differential centrifugation method that makes use of the higher density of negative electrostatic charges on luminal than on basolateral membranes to separate these fractions.

### Binding assay

Duplicate samples of <sup>125</sup>I-labeled toxin, either alone or in combination with varying amounts of

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unlabeled toxin, were incubated at the appropriate temperature with brush border membrane vesicles in a total volume of 100 ul of Tris buffer (Tris 10 mM, 150 mM NaCl, pH 7.4). All buffers contained 0.1 % BSA. The incubation temperature was 20 C. Ultrafiltration through Whatman GF/F glass fiber filters was used to separate bound from free toxin. Each filter was rapidly washed with 5 ml of ice-cold buffer (NaCl/P;- 0.1 % BSA). The radioactivity of the filter was measured in a gammacounter (1275 Minigamma, LKB). Binding data were analyzed using the LIGAND computer program. program calculates the bound concentration of ligand as a function of the total concentration of ligand, given the affinity (Ka or its inverse Kd = 1/Ka, the dissociation constant) and the total concentration of receptors or binding site concentration (R.).

### Determination of protein concentration

Protein concentrations of purified Bt2, Bt3, Bt73 and Bt15 toxins were calculated from the OD at 280 nm (measured with a Uvikon 810 P, Kontron Instruments spectrofotometer). The protein content of solutions of other toxins and of brush border membrane vesicles (BBMV) as measured according to Bradford (1976).

Binding of Bt2, Bt3 and Bt73 toxins to BBMV of Manduca sexta and Heliothis virescens: an example of 3 competitively binding Lepidopteran ICPs.

Bt2, Bt3 and Bt73 toxins are toxic to both Manduca sexta and Heliothis virescens: LC50 values for Manduca sexta are respectively 17.70, 20.20 and 9.00 ng/cm2; for Heliothis virescens the LC50's are 7.16, 90.00 and 1.60 ng/cm2.

Labelled toxin, either Bt3 (0.8 nM) or Bt2 (1.05 nM) or Bt73 (1.05 nM), was incubated with BBMV in a volume of 0.1 ml. BBMV protein concentrations were 100 ug/ml for M. sexta and for Bt2-H. virescens, for Bt3-H.

<u>virescens</u> 150 and for Bt73-<u>H</u>. <u>virescens</u> 50 ug/ml. The labelled toxin was combined with varying amounts of an unlabeled toxin (competitor). After a 30 min. incubation, bound and free toxins were separated through filtration.

Figs. 1-3 show the percentages binding respectively labelled Bt2, Bt3 and Bt73 toxins as a function of the concentration of competitor for Manduca sexta. Figs. 4-6 show these data for Heliothis virescens. The amount bound in the competitor is always taken as 100 % binding. Figs. 1-6 show the binding of  $^{125}I$ -labeled toxins to  $\underline{M}$ .  $\underline{\text{sexta}}$  (in Figs. 1, 2 and 3) and H. virescens (in Figs. 4, 5 and 6) brush border membrane vesicles. Vesicles were incubated with labeled toxin [in Figs. 1 and 4:  $^{125}I-$ Bt2-toxin (1.05nM); in Figs. 2 and 5:  $^{125}I-Bt3-toxin$ (0.8nM); in Figs. 3 and 6:  $^{125}I-Bt73-toxin$  (1.05nM)] in the presence of increasing concentrations of Bt2 toxin (\*), Bt3 toxin (●) or Bt73 toxin (▲). Binding is expressed as percentage of the amount bound upon incubation with labeled toxin alone. On M. sexta vesicles, these amounts were 1820, 601 and 2383 cpm, and on H. virescens vesicles 1775, 472 and 6608 cpm for <sup>125</sup>I-Bt2-, Bt3- and Bt73-toxin, respectively. specific binding was not substracted. were analyzed with the LIGAND computer program. Each point is the mean of a duplicate sample.

Figure 1: shows the binding of  $^{125}I$  Bt2 toxin to  $\underline{M}$ . sexta BBMV

Figure 2: shows the binding of  $^{125}I$  Bt3 toxin to  $\underline{M}$ . sexta BBMV

Figure 3: shows the binding of  $^{125}I$  Bt73 toxin to  $\underline{M}$ . sexta BBMV

Figure 4: shows the binding of  $^{125}$ I Bt2 toxin to  $\underline{\text{H}}$ .  $\underline{\text{virescens}}$  BBMV

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Figure 5: shows the binding of <sup>125</sup>I Bt3 toxin to H.virescens BBMV

Figure 6: shows the binding of <sup>125</sup>I Bt73 toxin to H.virescens BBMV

The conclusions from Figures 1-6 are that Bt2 and Bt3, Bt3 and Bt73, and Bt2 and Bt73 are competitively-binding ICP's both for <u>Manduca sexta</u> and for <u>Heliothis virescens</u>. Indeed Bt3 competes for the entire population of receptor sites of Bt2 in <u>Manduca sexta</u> (Fig.1): the % labelled Bt2 bound in the presence of 100 nM Bt3 is equal to the % Bt2 bound with 100 nM of Bt2 itself. The opposite is not true: in the presence of 100 nM Bt2 the % of labelled Bt3 is not reduced to the same level as with 100 nM of Bt3 (Fig.2).

A similar reasoning is followed to observe competitivity of other toxin combinations: Bt3 competes for the entire population of receptor sites of Bt73 (Fig. 3) in  $\underline{M}$ .  $\underline{sexta}$ ; the opposite is not true (Fig. 2); Bt2 and Bt73 compete for the entire population of each other's binding sites in  $\underline{M}$ .  $\underline{sexta}$  (Figs. 1 and 3).

In <u>Heliothis</u> <u>virescens</u>: Bt2 competes for the entire population of receptor sites of Bt3 (Fig. 5); Bt73 competes for the entire population of receptor sites of Bt3 (Fig. 5); Bt73 competes for the entire population of receptor sites of Bt2 (Fig. 4); but the opposite statements are not true (Figs. 4, 5 and 6).

The same data can be used in mathematical analysis (e.g., Scatchard analysis according to Scatchard, 1949; analysis with the LIGAND computer program according to Munson and Rodbard, 1980) to calculate the dissociation constant (Kd) of the toxin-receptor complex and the concentration of binding sites (Rt); the results of these calculations using the LIGAND computer program were the following:

Bt2- <u>M.sexta</u> :	Kd=0.4 nM	Rt=3.4	pmol/mg	vesicle
protein				
Bt3-M. sexta:	Kd=1.5 nM	Rt=9.8	pmol/mg	vesicle
protein				
Bt73- <u>M</u> . <u>sexta</u> :	Kd=0.6 nM	Rt=4.0	pmol/mg	vesicle
protein				
Bt2-H. virescens:	Kd=0.6 nM	Rt=9.7	pmol/mg	vesicle
protein				
Bt3-H. <u>virescens</u> :	Kd=1.2 nM	Rt=3.7	pmol/mg	vesicle
protein	•	•		
Bt73-H. <u>virescens</u> :	Kd=0.8 nM	Rt=19.5	pmol/mg	vesicle
protein				
				_

These data demonstrate the high affinity receptor binding of the toxins (Kds in the range of  $10^{-10}$  to  $10^{-9}$  M.

Binding of Bt2 and Bt14 toxins to BBMV of P. brassicae, Plutella xylostella and Phthorimaea opercullella: an example two non-competitively binding Lepidopteran ICPs

Bt2 and Bt14 toxins are toxic to <u>P. brassicae</u> (p.b.), <u>P. xylostella</u> (p.x.) and <u>P. operculella</u> (p.o.) as seen from the table below.

LC<sub>50</sub> of Toxins

	Bt2	Bt14
P.b.	1.3	2.0
P.x.	6.7	5.4
P.o.	4.20	0.8-4.0

 $LC_{50}$  values of solubilized purified Bt2 and Bt14 toxins for P.x. are expressed as ng protein spotted per cm<sup>2</sup> of artificial diet.  $LC_{50}$  values for P.b. are expressed as ug<sup>2</sup> toxin per ml solution into which leaf discs, fed to first instar Pb larvae, were dipped. For P.o.,  $LC_{50}$  values are expressed in ug/ml into which potato chips were dipped prior to feeding.

Labelled Bt2 toxin (1.05 nM) or Bt14 toxin (1.4 nM) was incubated with BBMV from P. brassicae (100 ug

protein/ml) in a volume of 0.1 ml in combination with varying amounts of unlabelled Bt2 or Bt14. After a 30 min. incubation period at 22°C, the bound and free toxins were separated.

Figures 7 and 8 show the binding of 125I-labeled toxins to P. brassicae brush border membrane vesicles. Vesicles were incubated with labeled toxin [in Fig. 7: 125I-Bt2-toxin (1.05nM); in Fig. 8: 125I-Bt14-toxin (1.4nM)] in the presence of increasing concentrations of Bt2 toxin (o) or Bt14 toxin (•). Binding is expressed as percentage of the amount bound upon incubation with labeled toxin alone. Non-specific binding was not substracted. Data were analyzed with the LIGAND computer program. Each point is the mean of a duplicate sample. Figure 7 shows the binding of labelled Bt2 toxin to P. brassicae BBMV, and Figure 8 shows the binding of labelled Bt14 toxin to  $\underline{p}$ . brassicae BBMV.

The competition data demonstrate the presence of high affinity binding sites both for Bt2 and Bt14, as well as the almost complete absence of competition of Bt14 for the Bt2 binding sites and of Bt14 for the Bt2 binding sites. This demonstrates that Bt2 and Bt14 are non-competitively binding toxins. Hence they are useful to prevent the development of <u>Pieris brassicae</u> resistance against <u>B. thuringiensis ICP's expressed in Brassica</u> sp.

Calculated Kd and Rt values were from these experiments were:

Bt2: Kd=2.8 nM, Rt=12.9 pmol/mg vesicle protein
Bt14: Kd=8.4 nM, Rt=21.4 pmol/mg vesicle protein.
Binding of Bt2 and Bt15 toxins to BBMV of M.sexta,
M.brassicae, P. xylostella and P.interpunctella: an
example of two non-competitively binding Lepidopteran
ICPs

Bt2 and Bt15 toxins are both toxic to  $\underline{\text{M.sexta}}$  (LC50's of 20 and 111 ng/cm2, respectively). They also show activity against  $\underline{\text{M. brassicae}}$ ,  $\underline{\text{P. xylostella}}$  and  $\underline{\text{P. interpunctella}}$ .

Labelled Bt2 (1.05 nM) or Bt15 (0.7 nM) was incubated with BBMV from M.sexta (100 ug protein/ ml) in a volume of 0.1 ml in combination with varying amounts of unlabelled Bt2 or Bt15. After a 30 min. incubation period at 22°C, the bound and free toxins were separated.

Figs. 9-10 show the binding of <sup>125</sup>I-labeled toxins to <u>M. sexta</u> brush border membrane vesicles. Vesicles were incubated with labeled toxin [in Fig. 9: <sup>125</sup>I-Bt2-toxin (1.05nM); in Fig. 10: <sup>125</sup>I-Bt15-toxin (0.7nM)] in the presence of increasing concentrations of Bt2-toxin (o) or Bt15-toxin (•). Binding is expressed as percentage of the amount bound upon incubation with labeled toxin alone. Non-specific binding was not substracted. Data were analyzed with the LIGAND computer program. Each point is the mean of a duplicate sample. Figure 9 shows the data for binding of labelled Bt2, and Figure 10 shows the binding of labelled Bt15.

The competition data demonstrate the presence of high affinity binding sites for both Bt2 and Bt15, as well as the complete absence of competition of Bt15 for the Bt2 binding sites and of Bt2 for the Bt15 binding sites. This demonstrates that Bt2 and Bt15 are non-competitively binding toxins. Hence the combination of Bt2 and Bt15 is useful to prevent the development of resistance of M.sexta against B. thuringiensis ICP's expressed in tobacco or other crops in which Manduca sp. are a pest. Calculated Kd and Rt values are:

Bt2: Kd=0.4 nM, Rt=3.4 pmol/mg vesicle protein

Bt15: Kd = 0.3 nM Kd2=2.9 nM, Rt1= 5.9 and Rt2=6.7 pmol/mg vesicle protein (2 distinct high affinity receptor sites are present).

Similar studies were performed for M. brassicae, S. littoralis and P. interpunctella. Although LD50, Kd and Rt values differed substantially, the essential observation that Bt2 and Bt15 are both toxic and are non-competitively binding toxins was confirmed in these three insect species. Thus, it is also a useful toxin combination to prevent resistance of M. brassicae to ICP's or to prevent resistance of Spodoptera species against ICP's expressed in any of the crop plants in which Spodoptera species are a pest.

Binding of Bt2 and Bt4 toxins to BBMV of M. sexta: an example of two non-competitively binding Lepidopteran ICPs

Both Bt2 and Bt4 toxins are toxic to Manduca sexta. LD50 values are 20 and 5.4. ng/cm2, respectively. No mutual competition of Bt2 for binding of labelled Bt4 and of Bt4 for binding of labelled Bt2 was observed, demonstrating that Bt2 and Bt4 are non-competitively binding toxins.

Binding of Bt15 and Bt18 toxins to BBMV of S. littoralis: an example of two non-competitively binding Lepidopteran ICPs

Both Bt15 and Bt18 toxins are toxic to S. littoralis. LD 50 values are 93 and 88 ng toxin/cm2, respectively. Labelled Bt15 (0.7 nM) or Bt18 (0.9 nM) was incubated with 100 ug of vesicle protein from S. littoralis in combination with varying amounts of unlabelled Bt15 or Bt18 toxin. After a incubation period, bound and free toxins separated. Binding data demonstrate high binding for both Bt15 and Bt18 to S. littoralis BBMV. As seen from Figures 11 and 12, the entire population of receptor sites of Bt15 was not saturable with Bt18, nor was the entire population of receptor sites of Bt18 saturable with Bt15.

Binding of Bt13 and Bt22 toxins to BBMV of L. decemlineata: an example of two non-competitively binding Coleopteran ICPs.

Both Bt13 and Bt22 toxins are toxic to L. decemlineata. LD 50 values are 0.8 and 1.1 ug toxin/ml respectively. Labelled Bt13 (1 nM) or Bt22 (0.7 nM) was incubated with 100 ug of vesicle protein/ml from S. littoralis in combination with varying amounts of unlabelled Bt13 or Bt22 toxin. After a 45 min. and incubation period, bound free toxins were separated. Binding data demonstrate high affinity binding for both Bt13 and Bt22 to S. littoralis BBMV. The entire population of receptor sites of Bt13 was not saturable with Bt22. Nor was the entire population of receptor sites of Bt22 saturable with Bt13.

Binding of Bt2 and Bt18 toxins to BBMV of M. sexta: an example of two non-competitively binding Lepidopteran ICPs.

Both Bt2 and Bt18 toxins are toxic to M. sexta, and LD 50 values are 20 to 73 ng toxin/cm² respectively. Labelled Bt2 (1.05nM) or Bt18 (0.7nM) was incubated with 100 ug/ml of vesicle protein from M. sexta in combination with varying amounts of unlabelled Bt2 or Bt18 toxin. After a 45 min. incubation period, bound and free toxins were separated. Binding data (Figs. 11-12) demonstrate high affinity binding for both Bt2 and Bt18 to M. sexta BBMV. The entire population of receptor sites of Bt2 was not saturable with Bt18. Nor was the entire population of receptor sites of Bt18 saturable with Bt2. Calculated Kd and Rt values are:

Bt2: Kd= 0.4 nM, Rt= 3.4 pmol/mg vesicle protein.

Bt18: Kd1= 0.04 nM, Rt1= 2.2 pmoles/mg vesicle protein and Kd2= 168nM Rt2= 194 pmoles/mg vesicle protein (2 distinct receptor sites for Bt18 are present).

A list of non-competitively binding anti-Lepidopteran ICP combinations and anti-Coleopteran ICP combinations is given below, together with their common target insect species in which non-competitivity has been demonstrated:

Bt2-Bt15 (Manduca sexta, Plutella xylostella, Pieris brassicae, Mamestra brassicae, Plodia interpunctella)

Bt2-Bt18 (Manduca sexta, Spodoptera littoralis)

Bt2-Bt14 (Pieris brassicae, Plutella xylostella,

Phthorimaea operculella)

Bt2-Bt4 (Manduca sexta)

Bt15-Bt18 (Manduca sexta, Spodoptera littoralis)

Bt14-Bt15 (Pieris brassicae)

Bt15-Bt4 (Manduca sexta, Spodoptera exiqua)

Bt18-Bt4 (Manduca sexta, Spodoptera littoralis)

Bt18-Bt14 (Pieris brassicae)

Bt18-Bt4 (Manduca sexta)

Bt13-Bt21 (Leptinotarsa decemlineata)

Bt13-Bt22 (Leptinotarsa decemlineata)

Bt21-Bt22 (Leptinotarsa decemlineata)

of course, this list of specific non-competitively binding ICP combinations for specific target insect pests is not exhaustive, and it is believed that other such ICP combinations, including combinations for yet-to-be discovered ICPs, will be found using a similar approach for any target insect species. Likewise, the foregoing list of target insect pests also is not exhaustive, and it is believed that other target insects pests (as well as the plants that are to be transformed to prevent their attack by such pests), against which the specific combinations of ICPs can be

used (e.g., the combination of the Bt2 and Bt14 ICPs in Brassica to prevent resistance of Pieris brassicae against the ICPs expressed in the plant), will be found using a similar approach.

# EXAMPLE 7: Selection for resistance of Manduca sexta (tobacco hornworm)

A selection experiment involves exposing a large number of larvae to a concentration of a toxin in a diet killing (e.g., 50-90 %) of the larvae. to toxin exposed surviving larvae are again concentrations killing a similar proportion of the larvae, and this process is continued for several generations. The sensitivity of the larvae to the toxin four generations investigated after each selection.

Selections for 20 generations of M. sexta were performed with Bt2 toxin alone, with Bt18 toxin alone and with a 1/4 (by weight) Bt2/Bt18 mixture. LC50 values of the reference strain for Bt2, Bt18 and the 1/4 Bt2/Bt18 mixture respectively were the following: 20 ng/cm2, 73 ng/cm2 and 62 ng/cm2 of diet.

Selection was initiated at concentrations killing around 75 % of the larvae. After 4 generations of selection, survival increased in both the Bt2 and the Bt18 selection to around 70 %, no such increase was observed in the selection with the combination of Bt2 and Bt18. Dosages were again increased to calculated LC75 values. This was repeated every 4 generations. The selection process was thus continued to the 20th generation. Final results were the following (LC50 of the 20th generation):

- Bt2 selection: LC50 was 6400 ug/g (320 times decreased sensitivity)
- Bt18 selection: LC50 was 15100 ug/g (207 times decreased sensitivity)

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- Bt2/Bt18 selection: LC50 was 181 ug/g (3 times decreased sensitivity).

Thus the decrease in sensitivity was about 100 times slower in the combined selection experiment.

Receptor binding in the three selected M. sexta strains was investigated with Bt2 and Bt18 and compared to those of the reference M. sexta strain (non-selected strain). Binding characteristics of the reference strain for the Bt2 and BT18 toxins were:

Bt2: Kd = 0.4 nM, Rt=3.4 pmol/mg vesicle protein
Bt18: Kd1=0.04 nM, Rt1=2.2 pmoles/mg vesicle protein
and Kd2=168nM, Rt2=194 pmoles/mg vesicle protein (2
distinct receptor sites for Bt18 are present).

Figures 11 and 12 show the binding of <sup>125</sup>I-labeled toxins to <u>M</u>. <u>sexta</u> brush border membrane vesicle. Vesicles were incubated with labeled toxin [in Fig. 11: <sup>125</sup>I-Bt2-toxin (1.05nM); in Fig. 12: <sup>125</sup>I-Bt18-toxin (0.7nM)] in the presence of increasing concentrations of Bt2-toxin (0) or Bt18-toxin (•). Binding is expressed as percentage of the amount bound upon incubation with labeled toxin alone. Non-specific binding was not substracted. Data were analyzed with the LIGAND computer program. Each point is the mean of a duplicate sample.

The Bt2 selected strain showed no detectable high affinity binding of Bt2 whereas its Bt18 binding characteristics remained close to the reference strain. (Bt18: Kd1=0.03 nM, Rt1=2.8 pmoles/mg vesicle protein and Kd2=199nM, Rt2=109 pmoles/mg vesicle protein; 2 distinct receptor sites for Bt18 are still present).

The Bt18 selected strain lost the high affinity receptor site for Bt18. The lower affinity site for Bt18 was still present in lower concentration than in the reference strain (Kd=189 nM, Rt=43 nM). Bt2 binding site concentration increased markedly compared to the

reference strain (Kd=0.4 nM, Rt=20.8 pmoles/mg vesicle protein). This strain had a Bt2 sensitivity of  $LC_{50}$ =4 ng/cm<sup>2</sup>. Thus, its sensitivity for Bt2 had increased as compared to the reference strain ( $LC_{50}$ =20 ng/cm<sup>2</sup>).

The Bt2/Bt18 selected strain showed a slight but statistically non-significant decrease in Bt18 binding site concentration. (Bt2: Kd = 0.4 nM, Rt=3.4 pmol/mg vesicle protein, Bt18 : Kd1=0.04 nM, Rt1=1.0 pmoles/mg vesicle protein and Kd2=168nM, Rt2=194 vesicle protein; 2 distinct receptor sites for Bt18 are present). These data demonstrate that, in the two selection lines where resistance occurred, the mechanism was situated at the receptor level. Changes in receptor site are shown to be the most likely mechanism of resistance to B. thuringiensis ICPs.

EXAMPLE 8: Mechanism of resistance of the diamondback moth to the microbial insecticide Bacillus thuringiensis.

The mechanism of development of insect resistance to ICPs has been investigated in a P. xylostella strain ("PxR"). This insect strain has developed a high level of resistance in the field against Dipel. Crystals of Dipel preparations contain a mixture of ICPs such as Bt3, Bt2 and Bt73 ICPs; in Example 6, it has been shown that these toxins are competitively binding ICPs.

Resistance to Dipel was confirmed by the toxicity data for the sensitive strain ("PxS") and for the Dipel-resistant strain ("PxR"). High levels of resistance are also observed for the Bt2 protoxin and toxin as shown in the following table:

LC<sub>50</sub> of Strains

	PxS	PxR	
Bt2	6.7	> 1350	
Bt15	132.6	120.4	

 $LC_{50}$  data are expressed as ng protein spotted per cm<sup>2</sup> of artificial diet.

However, insect toxicity data show that there is no resistance to the Bt15 protoxin and Bt15 toxin; this ICP is not present in Dipel crystals. To investigate whether a change in toxin-membrane binding was responsible for resistance, receptor binding studies were performed with <sup>125</sup>I-labeled Bt2 toxin and Bt15 toxin, with BBMV derived from larvae midguts of the PxR and PxS strains. The results are summarized in Table 1, below.

Table 1. Binding characteristics of Bt2 and Bt15 toxins to brush border membrane vesicles from sensitive and resistant P. xylostella.

ICP	strain	Kd (nM)	Rt (pmol/
			mg protein)
Bt2 toxin	PxS	8.1	1.6
	PxR	no binding	detectable
Bt15 toxin	PxS	1.9	4.2
	PxR	3.7	5.8

Table 1 shows that there was high-affinity saturable binding of the Bt2 toxin to midgut membranes of the PxS strain, but the PxR strain showed no detectable level of Bt2 toxin binding. With the Bt15 toxin, there was significant binding to BBMW of both the PxR and PxS strains, and values are not significantly different for the two strains.

These data show that resistance in <u>P. xylostella</u> is due to an alteration in toxin-membrane binding. Resistance to the Bt2 toxin and the sensitivity toward the Bt15 toxin of the PxR strain is reflected by the binding characteristics shown in Table 1.

Hence, when different non-competitively binding ICPs (i.e., Bt2 and Bt15) are available with activity against the same insect species (e.g., P. xylostella),

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resistance to one ICP(Bt2) does not imply resistance against other ICPs (such as Bt15). Thus, ICPs with different binding properties can be used in combination to delay development of insect resistance to ICPs.

EXAMPLE 9: Separate transfer of two ICP genes within individual transcriptional units to the genome of plant cells

Two procedures are envisaged for obtaining the combined expression of two ICP genes, such as the <u>bt2</u> and <u>bt15</u> genes in transgenic plants, such as tomato plants. These procedures are based on the transfer of two chimeric ICP genes, not linked within the same DNA fragment, to the genome of a plant of interest.

A first procedure is based on sequential transformation steps in which a plant, already transformed with a first chimeric ICP retransformed in order to introduce a second ICP gene. sequential transformation makes use two different selectable marker genes, such as the resistance genes for kanamycin ("km") and phosphinotricin acetyl transferase ("PPT"), which confers resistance to phoshinotricin. The use of both these selectable markers has been described in De Block et al. (1987).

The second procedure is based on the cotransformation of two chimeric ICP genes on different plasmids in a single step. The integration of both ICP genes can be selected by making use of the two selectable markers conferring resistance to Km and PPT, linked with the respective ICP genes.

For either procedure, a Ti-plasmid vector is used for Agrobacterium-mediated transformation of each chimeric ICP gene into plant cells.

Plasmid pGSH163, described in EP 0193259, contains the following chimeric genes between the T-DNA border

repeats: a gene fragment encoding the toxin part of the bt2 gene under the control of the TR2' promoter and the neo gene under control of the TR1' promoter. The 3' ends of the T-DNA gene 7 and octopine synthase respectively provide information for the 3' end formation of transcripts.

encoding the toxin of the Bt15 ICP under the control of the TR2' promoter, was constructed in the following way (Figure 15). pOH50 consists of pUC18 with the whole bt15 gene under the control of the lac promoter. A HindIII-BglII fragment was cloned in pMa5-8 yielding pJB3. By site-directed mutagenesis, a NcoI site was created at the initiation codon to yield pVE29. A fragment containing the truncated gene fragment of the bt15 gene, with a translational stop codon, was obtained by isolation of BclI-ClaI from pOH50 and cloning in pLK91, yielding pHW38. The whole toxin gene fragment was reconstructed under the control of the tac promoter, yielding pVE35, by ligation of a ClaI-PstI fragment from pHW38, a NcoI-ClaI fragment from pVE29 and a NcoI-PstI fragment from pOH48. A truncated bt15 gene fragment with a NcoI site at the initiation codon was obtained from pVE35 as a 1980 NcoI-BamHI fragment and cloned in pGSJ141, digested with ClaI and BamHI. pGSJ141 has been described in EPA 88402115.5. Ligation of the filled ClaI site to the filled NcoI site yielded a chimeric TR2' - truncated bt15 - 3'g7 construct (pTVE47). As a selectable marker in this plasmid, the bar gene encoding phosphinothricin acetyl transferase and conferring resistance to PPT was used. A chimeric bar gene containing the bar gene under the control of the 35S promoter and followed by the 3' end of the octopine synthase was introduced in pTVE47. From pDE110, a 35S-bar-3'ocs fragment was obtained as a

StuI-HindIII fragment and was cloned in pTVE47 digested with PstI and HindIII. This yielded the plasmid pTHW88 (Figure 15) which contains the truncated <u>bt15</u> gene under the control of the TR2' promoter and the bar gene under the control of the 35S promoter between the T-DNA border repeats. Plasmid pGSH163 is cointegration type Ti-plasmid vector, whereas pTHW88 is a binary type Ti-plasmid vector as described in EPA 0193259.

Both plasmids were mobilized in the A. tumefaciens strain C58C1Rif (pGV2260) according to Deblaere et al. (1988). In the sequential transformation procedure, tomato was transformed according to De Block et al. (1987) with the A. tumefaciens strain C58C1Rif carrying pGS1163 resulting from the cointegration of pGSH163 and pGV2260. Individual transformants were selected for kanamycin resistance, and regenerated plants were characterized for expression of the truncated bt2 gene according to Vaeck et al. (1987). One representative transformant was subsequently retransformed with the  $\underline{A}$ . tumefaciens strain C58C1Rif (pGV2260 and pTHW88), and transformants were selected for PPT resistance. Using cotransformation procedure, the Agrobacteria strains, carrying the cointegrate vector pGS1163 and the binary vector pTHW88, were used for transformation of tomato. Individual plants selected for resistance to Km and PPT.

Schematically shown in Fig. 15 are:

- b) construction of pVE35: <a href="https://doi.org/bt/b/b/b/b/b/b/b/b/b/5/">bt15</a> C-terminal truncated gene fragment under control of the tac promoter.

c) construction of pTHW88: binary T-DNA vector with a chimeric <u>bt15</u> gene and a chimeric <u>bar</u> gene within the T-DNA border repeats.

In both cases, co-expression of the two ICP genes in the individual transformants was evaluated by insect toxicity tests as described in EP 0193259 and by biochemical means. Specific RNA probes allowed the quantitive analysis of the transcript levels; antibodies cross-reacting monoclonal respective gene products allowed the quantitative analysis of the respective gene products in ELISA tests (EP 0193259); and specific DNA probes allowed the characterization of the genomic integrations of the bt2 and bt15 genes in the transformants. It was that the transformed tomato plants found simultaneously expressed both the bt2 gene (8.1 ng/mg) and the bt15 gene (7.6 ng/mg) as measured by ELISA, which would prevent or delay development of resistance of M. sexta to the insecticidal effects of the Bt2 and Bt15 toxins, being expressed.

These procedures also could be applied when one or both ICP genes are part of a hybrid gene. For example, the same strategy as described above could be followed with the plasmid vectors pGSH152, containing a chimeric truncated <a href="https://doi.org/10.1001/journal.org/10.1001/journa

EXAMPLE 10: Separate transfer of two ICP genes to the nuclear genome of separate plants in independent transformation events and subsequent combination in a single plant through crossing.

Tobacco plants have been transformed with either the <a href="https://doi.org/10.1001/journal.com/bt18">bt18</a> gene or the <a href="https://doi.org/10.1001/journal.com/bt18">bt18</a> gene or the <a href="https://doi.org/10.1001/journal.com/bt18">bt18</a> gene or the <a href="https://doi.org/10.1001/journal.com/bt18">bt18</a> gene by applying the same cloning strategies as described in EP 0358557 and EP

193259, respectively. For both genes, the plants were transformed with plant expression vectors containing either the truncated <u>bt18</u> or <u>bt15</u> gene, which just encode the Bt18 or Bt15 toxin, respectively.

The mortality rate of <u>Spodoptera littoralis</u> larvae feeding on the transformed plants is significantly higher than the mortality rate of larvae fed on untransformed plants.

The <u>bt18</u>-transformed plant, which is homozygous for the <u>bt18</u> gene, is then crossed with the <u>bt15</u> - transformed plant, which is homozygous for the <u>bt15</u> gene. After selfing, a plant homozygous for both genes is obtained.

The resulting tobacco plants, expressing both the <u>bt18</u> and <u>bt15</u> genes, delay significantly development of resistance by <u>S. littoralis</u> to either the Bt18 or Bt15 toxin expressed by the plants.

## EXAMPLE 11: Transfer of two chimeric ICP genes linked within the same DNA to the genome of plant cells

The strategy used is based on the organization of two independent chimeric ICP genes between the T-DNA border repeats of a single vector. Binding studies indicated that the Bt2 and Bt14 toxins are two non-competitively binding ICPs with insecticidal activity towards Pieris brassicae. For expression in plants, both the bt2 and bt14 genes can be coexpressed to prevent insect resistance development. For the design of a plasmid vector with each ICP gene under the control of a separate promoter, possibilities can be envisaged: 1) three chimeric constructs carrying the truncated bt2 and bt14 genes and a selectable marker, respectively; or 2) a hybrid of a truncated gene fragment (bt2 or bt14) and the neo gene can be used in combination with a truncated bt14 or bt2 gene.

This Example describes the construction of the vector pTHW94 for plant transformations carrying the following chimeric ICP genes between the T-DNA border repeats: a truncated <u>bt2</u> gene fragment under the control of the TR2' promoter and a hybrid truncated <u>bt14-neo</u> gene under the control of the TR1' promoter. The 3' end of the T-DNA gene 7 and octopine synthase, respectively, provide information for proper 3' end formation. pTHW94 has been deposited at the DSM under accession no. 5514 on August 28, 1989.

Schematically shown in Fig. 16 are the:

- b) construction of pHW67: reconstruction of the  $\frac{bt14}{control}$  gene under the control of the tac promoter.
- c) construction of pHW71: construction of a hybrid truncated <a href="https://doi.org/block.org/bh/b/-100/bh/-
- d) construction of pTHW94: binary T-DNA vector with a chimeric <u>bt14</u> gene and a chimeric <u>bt2</u> gene within the T-DNA border repeats.

The pTHW94 vector is mobilized into the Agrobacterium strain C58C1Rif (pMP90) which is used to transform Brassica napus according to the procedure described by De Block et al. (1989). Transformants are selected on Km, and regenerated plants are found to express both ICP gene products in insect toxicity tests and biochemical tests.

## EXAMPLE 12: Expression of two ICP genes in a hybrid construct

In order to obtain a combined and simultaneous expression of two ICP genes, truncated gene fragments encoding the toxic parts of two different ICPs can be fused in a proper reading frame and placed, as a hybrid gene, under the control of the same promoter in a chimaeric gene construct. Toxic cores from certain ICPs can be liberated from their protoxins by protease activation at the N- and/or C- terminal end. Thus, hybrid genes can be designed with one or more regions encoding protease cleavage site(s) at the fusion point(s) of two or more ICP genes.

The simultaneous co-expression of the bt2 and bt14 genes is obtained by constructing a hybrid gene composed of a truncated bt14 gene fragment fused to a truncated bt2 gene fragment. Schematically shown in Figure 17 is the construction of such a hybrid bt2-bt14 gene with a C-terminal bt2 gene fragment (bt860) encoding the toxic core of the Bt2 protoxin in frame with a C-terminal truncated bt14 gene fragment encoding the toxic core of the protoxin. The BclI site in the bt2 gene, localized downstream of the trypsin cleavage site, is fused in frame with the NcoI site introduced at the N-terminal end of the truncated bt14 gene fragment. To this end, the plasmids pLBKm860 (EP 0193259) and pHW67 are used. pLBKm860 contains a hybrid bt2-neo gene under control of the lambda Pi promoter. The bt2 gene moiety in the hybrid gene is a C-terminal truncated bt2 gene fragment, indicated as bt860 (in Fig. 17) (see also Vaeck et al, 1987). The construction of pHW67 is described in Fig. 16. pHW67 contains a Cterminal truncated bt14 gene fragment (bt14tox) with NcoI site at the ATG initiation

translation stop codon located at the BclI site of the intact <u>bt14</u> gene and a BamHI site downstream of the whole gene fragment. To fuse both gene fragments in the proper reading frame, the BclI and NcoI ends of the respective plasmids are treated with Klenow DNA polymerase and S1 nuclease as indicated in Figure 16. The resulting plasmid pJB100 contains the hybrid <u>bt860-bt14tox</u> gene under control of the lambda P<sub>L</sub> promoter and directs the expression in <u>E. coli</u> of a fusion protein with the expected mobility on SDS-PAGE.

Crude extracts of the <u>E</u>. <u>coli</u> strain show the toxicity of the fusion protein, expressed by the strain, against <u>P</u>. <u>brassicae</u>. It has also been confirmed by N-terminal amino acid sequence analyses of the fusion protein produced by the <u>E</u>. <u>coli</u> strain that the N-terminal amino acids from the Bt14 protoxin are processed upon activation. The <u>bt2-bt14</u> hybrid gene product has thus two potential protease cleavage sites.

Subsequently, this hybrid gene is inserted into a vector for plant transformations and placed under control of a suitable promoter and transferred to the genome of brassica (EP 0193259) where both the <u>bt2</u> and <u>bt14</u> genes are expressed in insect toxicity tests.

Table 2

Gene	Bt strain	Host range	amino acids - encoded		Disclosure of nucleotide sequence
					·
bt3	HD-1 kurstaki	L	1176	133.2	Schnepf et al.,1985
bt2	berliner 1715	L	1155	131	Höfte et al.,1986
bt73	HD-73	L	1178	133.3	Adang et al, 1985
bt14	entomocidus HD-110	Ĺ	1207	138	Brizzard and Whiteley,
	• :				1988
bt15	entomocidus HD-110	L	1189	134.8	Fig. 14
bt4	HD-68 aizawai	L	1165	132.5	Fig. 13
bt18	darmstadiensis HD-146	L	1171	133	EP appln. 88402241.9
bt13	Bt51,D5M4288 22/10/87	С	644	73.1	EP appln. 88402115.5
bt21	DSM 5131, 19/1/89	С	651	74.2	EP appln. 89400428.2
bt22	BtPGS1245, DSM 5132, 19/1/89	C	1138	129	EP appln. 8940028.2
P2	HD-263	L/D	633	70.9	Donovan et al, 1988
Cry B2	HD-1	Ĺ	633	70.8	Widner and Whiteley, 1989

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\*

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#### **CLAIMS**

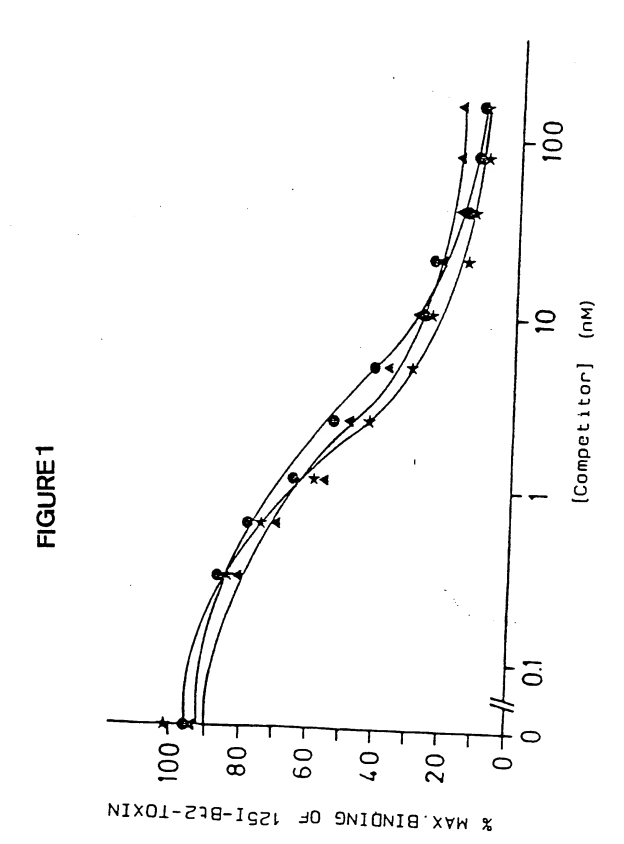
- 1. A cell of a plant, characterized by: at least two <u>B</u>. <u>thuringiensis</u> ICP genes stably inserted into the genome of said plant; each of said genes encoding a different non-competitively binding ICP for an insect species; whereby at least two different ICPs can be produced by said cell which do not bind competitively to the brush border membrane of the columnar midgut epithelial cell of said insect species.
- 2. The cell of claim 1 wherein at least one marker gene, encoding a protein or polypeptide which renders said cell easily distinguishable from cells which do not contain said protein or polypeptide, is in the same genetic locus as at least one of said ICP genes.
- 3. The cell of claim 1 or 2, wherein each of said ICP genes is under the control of a separate promoter capable of directing gene expression in said cell and is provided with a separate signal for 3' end formation and within a same transcriptional unit.
- 4. The cell of claim 2 or 3, in which said marker DNA is under the control of a separate promoter capable of directing gene expression in said plant cell and is provided with a signal for 3' end formation within a same transcriptional unit.
- 5. The cell of claim 1 or 2, wherein said ICP genes are within a same transcriptional unit and under the control of a single promoter.
- 6. The cell of claim 5, wherein said marker gene is fused with at least one of said ICP genes and is within said same transcriptional unit and under the control of said promoter.

- 7. The cell of claim 5 or 6, wherein a DNA fragment, encoding a protease-sensitive or cleavable amino acid sequence, is in said same transcriptional unit as said ICP genes and intercalated in frame between said ICP genes.
- 8. The cell of claim 5 or 6, wherein said ICP genes are combined in a dicistronic unit comprising an intergenic DNA sequence which allows reinitiation of translation and is in said same transcriptional unit as said ICP genes and intercalated between said ICP genes.
- 9. The cell of anyone of claims 1 to 8, wherein said ICP genes are genes encoding insecticidal proteins having activity against Lepidoptera species and are particularly the following genes: bt2 and/or bt73 and/or bt4 and/or bt14 and/or bt15 and/or bt18.
- 10. The cell of any of claims 1 to 8, wherein said ICP genes are genes encoding insecticidal proteins having activity against a Coleoptera species and are particularly the following genes: bt13 and/or bt21 and/or bt22.
- 11. The cell of any of claims 2 to 10 wherein said marker DNA is: an herbicide resistance gene, particularly a <u>sfr</u> or <u>sfrv</u> gene; a gene encoding a modified target enzyme for a herbicide having a lower affinity for the herbicide, particularly a modified 5-EPSP as a target for glyphosate or a modified glutamine synthetase as a target for a GS inhibitor; or an antibiotic resistance gene, particularly NPTII.
- 12. The cell of any of claims 3 to 6, wherein said promoter is: a constitutive promoter, particularly a 35S promoter or a 35S3 promoter; a PNOS promoter; a POCS promoter; a wound-inducible

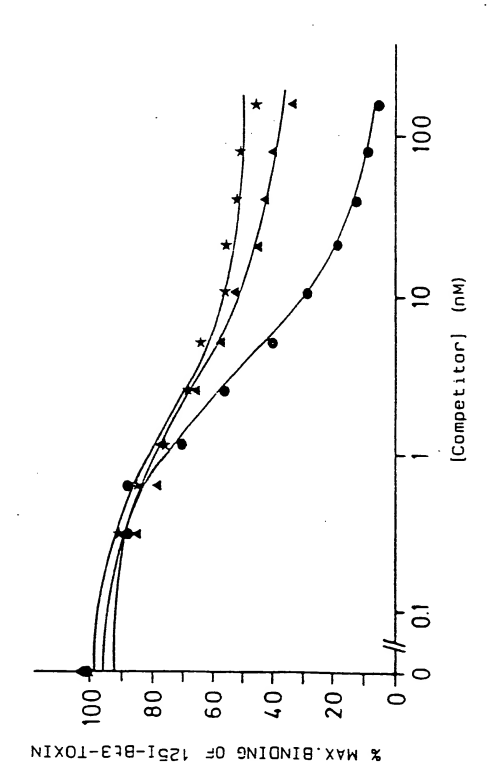
promoter, particularly a TR1' or TR2' promoter; a promoter which directs gene expression selectively in plant tissue having photosynthetic activity, particularly a SSU promoter; or a tissue-specific promoter, particularly a tuber-specific promoter, a stem-specific promoter or a seed-specific promoter.

- 13. A vector suitable for transforming a cell of a plant, particularly a plant capable of being infected with <u>Agrobacterium</u>, comprising said ICP genes of any of claims 1 to 12.
- 14. A process for producing a plant having improved insect resistance and having said ICP genes of anyone of claims 1 to 12 stably integrated into the nuclear genome of their cells, characterized by the non-biological steps of transforming a cell of said plant by introducing said ICP genes into the nuclear genome of said cell and regenerating said plant and reproduction material from said cell.
- 15. A plant cell culture, consisting of the plant cells of anyone of claims 1 to 12.
- 16. A plant, consisting of the plant cells of anyone of claims 1 to 12.
- 17. Brassica, tomato, potato, tobacco, cotton or lettuce consisting of the plant cells of anyone of claims 1 to 12, wherein said ICP genes comprise one of the following pairs of genes: bt2 and bt18 or bt73 and bt15 or bt2 and bt18 or bt2 and bt14 or bt2 and bt15 or bt14 and bt15 or bt4 and bt15 or bt13 and bt21 or bt21 and bt22 or bt13 and bt22.
- 18. The cell of anyone of claims 1-12, made by a process as described hereinabove.

19. A method for rendering a plant resistant to an insect species by transforming the plant with said ICP genes of anyone of claims 1-12.







က FIGURE

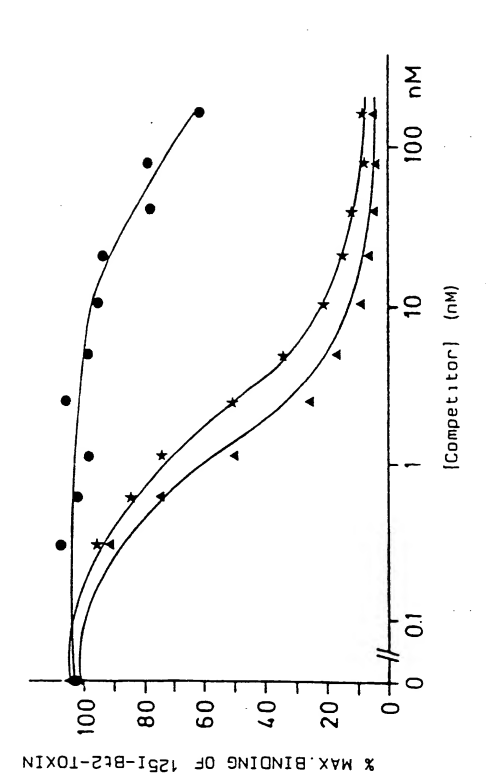


FIGURE 4

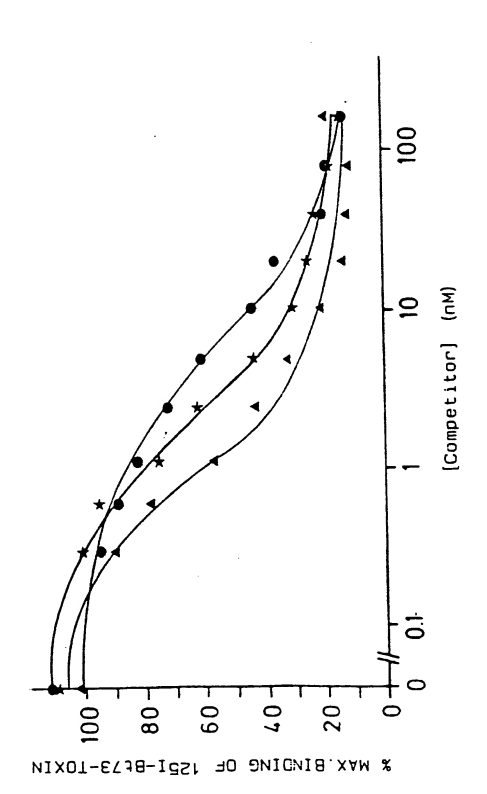


FIGURE 5

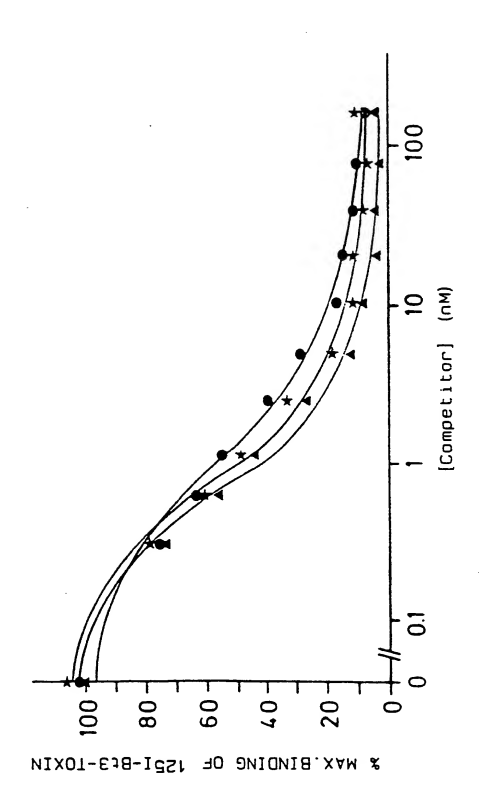
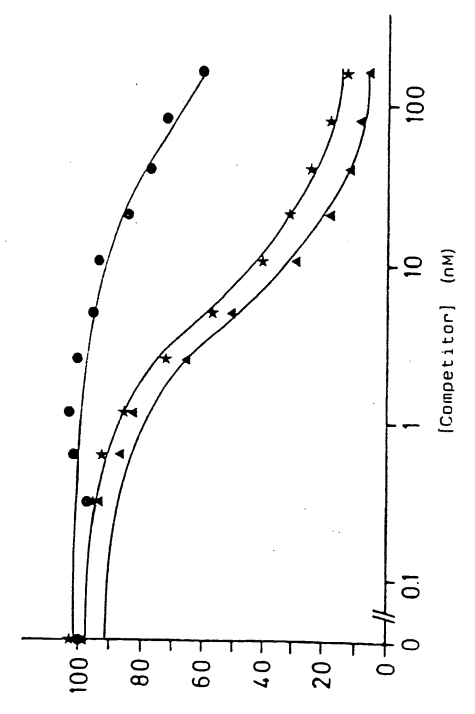
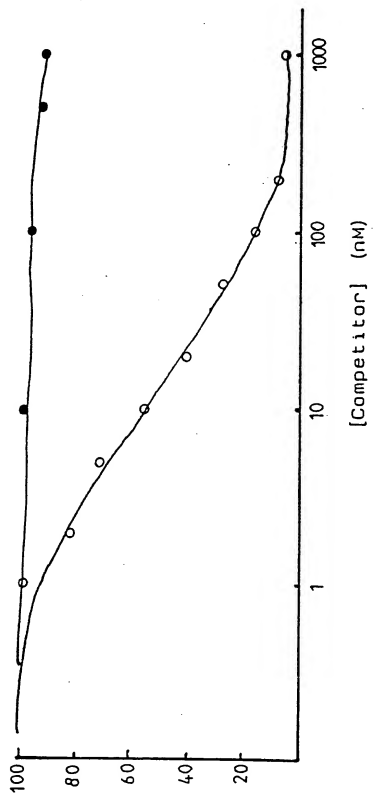


FIGURE 6

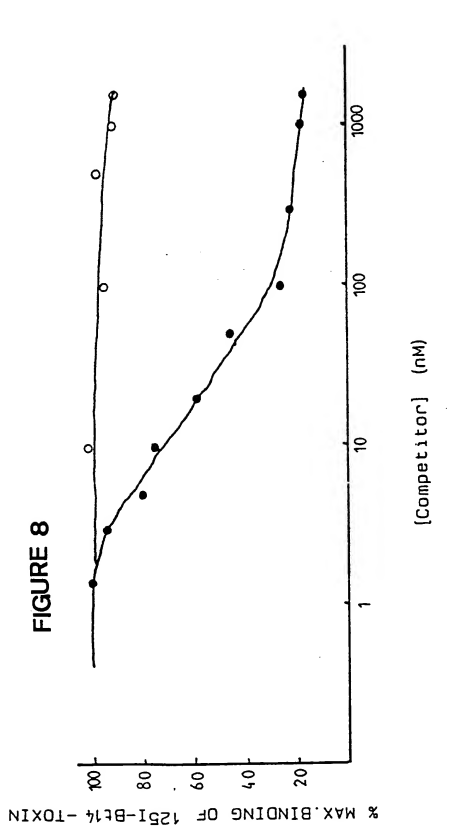


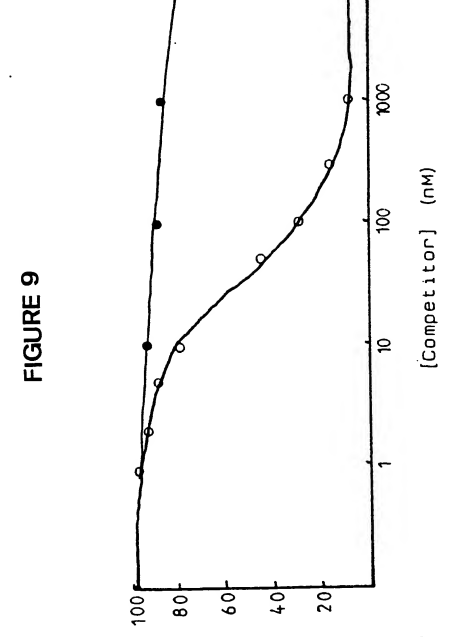
% MAX.BINDING OF 1251-BL73-TOXIN





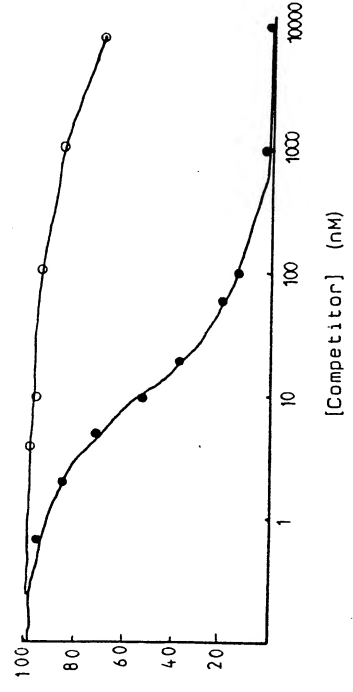
% MAX.BINDING OF 1251-BL2-TOXIN





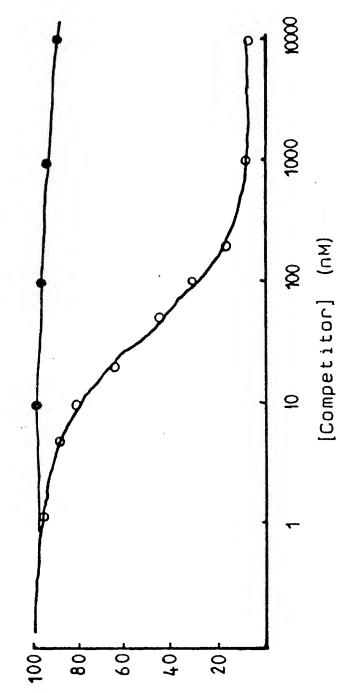
% MAX.BINDING OF 1251-BL2-TOXIN

FIGURE 10



% MAX.BINDING OF 1251-BL15 -TOXIN





% MAX.BINDING OF 1251-Bt2-TOXIN

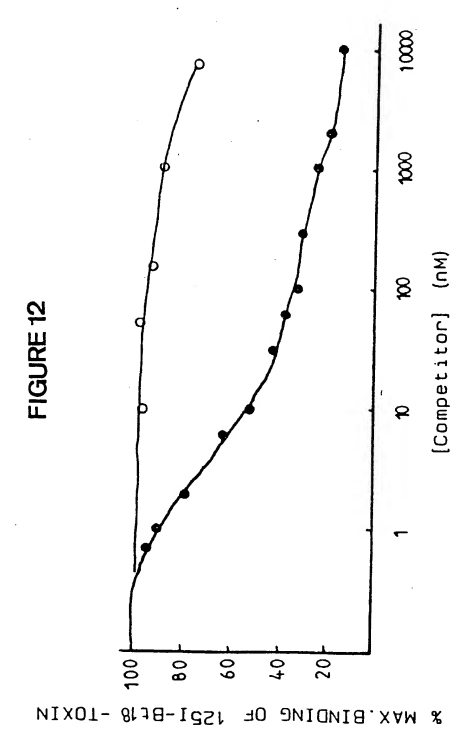


Figure 13

GGATCTGTTT TAATATAAGG GATTTGTGCC CTTCTCGTTA TATTCTTTTA TTAGCCCCAA AAACTAGTGC AACTAAATAT TTTTATAATT ACACTGATTA 110 120 130 140 AATACTTTAT TTTTGGGAGT AAGATTTATG CTGAAATGTA ATAAAATTCG 170 180 TTCCATTTTC TGTATTTTCT CATAAAATGT TTCATATGCT TTAAATTGTA GTAAAGAAAA ACAGTACAAA CTTAAAAAGGA CTTTAGTAAT TTAATAAAAA

260 269 278 287

AAGGGGATAG TTT ATG GAA ATA AAT AAT CAA AAC CAA TGT

MET Glu Ile Asn Asn Gln Asn Gln Cys

Figure 13 (Cont. 1)

GTG CCT TAC AAT TGT TTA AGT AAT CCT AAG GAG ATA ATA Val Pro Tyr Asn Cys Leu Ser Asn Pro Lys Glu Ile Ile TTA GGC GAG GAA AGG CTA GAA ACA GGG AAT ACT GTA GCA Leu Gly Glu Glu Arg Leu Glu Thr Gly Asn Thr Val Ala GAC ATT TCA TTA GGG CTT ATT AAT TTT CTA TAT TCT AAT Asp Ile Ser Leu Gly Leu Ile Asn Phe Leu Tyr Ser Asn TTT GTA CCA GGA GGA GGA TTT ATA GTA GGT TTA CTA GAA Phe Val Pro Gly Gly Gly Phe Ile Val Gly Leu Leu Glu TTA ATA TGG GGA TTT ATA GGG CCT TCG CAA TGG GAT ATT Leu Ile Trp Gly Phe Ile Gly Pro Ser Gln Trp Asp Ile TTT TTA GCT CAA ATT GAG CAA TTG ATT AGT CAA AGA ATA Phe Leu Ala Gln Ile Glu Gln Leu Ile Ser Gln Arg Ile Figure 13 (Cont. 2)

	530			539			548			557		
							GCA					
							Ala					
							GTC					
Gly	Leu	Ser	Asn	Leu	Tyr	Lys	Val	Tyr	Val	Arg	Ala	Phe
		(11			600			600				
	~~~											
							ACT					
Ser	Asp	Trp	Glu	Lys	Asp	Pro	Thr	Asn	Pro	Ala	Leu	Arg
	617			656			(()			COL		
011	04/	1 770	000	000	<b></b>		665			0/4		
							AAT					
Glu	Glu	MET	Arg	Ile	Gln	Phe	Asn	Asp	MET	Asn	Ser	Ala
602			602			701			710	r		710
							TTT					
ьeu	116	TNI	ATA	116	Pro	Leu	Phe	Arg	Val	Gln	Asn	Tyr
		728			737			746			755	
CDD	乙中中						TAT					
_												
C111	Val	Ala	LAU	I All	$C \sim \sim$	Wal	Mers.	17.1	(ነ ∽	11.	Ala	1

Figure 13 (Cont. 3)

	764			773	•		782			791		
TTA												GGA
												Gly
800			809			818			827			836
					GAT							
					Asp							
		845			854			863			872	
TAT	AGT	GAT	CTG	ACT	AGC	CTT	ATT	CAT	GTT	TAT	ACT	AAC
Tyr	Ser	Asp	Leu	Thr	Ser	Leu	Ile	His	Val	Tyr	Thr	Asn
	881			890			899			908		
CAT	TGT	GTG	GAT	ACG	TAT	AAT	CAG	GGA	TTA	AGG	CGT	TTG
His	Cys	Val	Asp	Thr	Tyr	Asn	Gln	Gly	Leu	Arg	Arg	Leu
917			926			935			944			953
GAA	GGT	CGT	TTT	CTT	AGC	GAT	TGG	ATT	GTA	TAT	AAT	CGT
Glu	Gly	Arg	Phe	Leu	Ser	Asp	Trp	Ile	Val	Tyr	Asn	Arg
		962			971			980			989	
TTC	CGG	AGA	CAA	TTG	ACA	ATT	TCA		TTA	GAT		GTT
Phe	Arg	Arg	Gln	Leu	Thr	Ile	Ser	Val	Leu	Asp	Ile	Val
										_		

Figure 13 (Cont. 4)

998 1007 1016 1025 GCG TTT TTT CCA AAT TAT GAT ATT AGA ACA TAT CCA ATT Ala Phe Phe Pro Asn Tyr Asp Ile Arg Thr Tyr Pro Ile

1034 1043 1052 1061 1070 CAA ACA GCT ACT CAG CTA ACG AGG GAA GTC TAT CTG GAT Gln Thr Ala Thr Gln Leu Thr Arg Glu Val Tyr Leu Asp

1079 1088 1097 1106
TTA CCT TTT ATT AAT GAA AAT CTT TCT CCT GCA GCA AGC
Leu Pro Phe Ile Asn Glu Asn Leu Ser Pro Ala Ala Ser

1115 1124 1133 1142
TAT CCA ACC TTT TCA GCT GCT GAA AGT GCT ATA ATT AGA
Tyr Pro Thr Phe Ser Ala Ala Glu Ser Ala Ile Ile Arg

1151 1160 1169 1178 1187

AGT CCT CAT TTA GTA GAC TTT TTA AAT AGC TTT ACC ATT

Ser Pro His Leu Val Asp Phe Leu Asn Ser Phe Thr Ile

1196 1205 1214 1223
TAT ACA GAT AGT CTG GCA CGT TAT GCA TAT TGG GGA GGG
Tyr Thr Asp Ser Leu Ala Arg Tyr Ala Tyr Trp Gly Gly

Figure 13 (Cont. 5)

1232 1241 1250 1259
CAC TTG GTA AAT TCT TTC CGC ACA GGA ACC ACT ACT AAT
His Leu Val Asn Ser Phe Arg Thr Gly Thr Thr Thr Asn

1268 1277 1286 1295 1304
TTG ATA AGA TCC CCT TTA TAT GGA AGG GAA GGA AAT ACA
Leu Ile Arg Ser Pro Leu Tyr Gly Arg Glu Gly Asn Thr

1313 1322 1331 1340
GAG CGC CCC GTA ACT ATT ACC GCA TCA CCT AGC GTA CCA
Glu Arg Pro Val Thr Ile Thr Ala Ser Pro Ser Val Pro

1349 1358 1367 1376
ATA TTT AGA ACA CTT TCA TAT ATT ACA GGC CTT GAC AAT
Ile Phe Arg Thr Leu Ser Tyr Ile Thr Gly Leu Asp Asn

1385 1394 1403 1412 1421 TCA AAT CCT GTA GCT GGA ATC GAG GGA GTG GAA TTC CAA Ser Asn Pro Val Ala Gly Ile Glu Gly Val Glu Phe Gln

1430 1439 1448 1457

AAT ACT ATA AGT AGA AGT ATC TAT CGT AAA AGC GGT CCA

Asn Thr Ile Ser Arg Ser Ile Tyr Arg Lys Ser Gly Pro

Figure 13 (Cont. 6)

1466 1475 1484 1493
ATA GAT TCT TTT AGT GAA TTA CCA CCT CAA GAT GCC AGC
Ile Asp Ser Phe Ser Glu Leu Pro Pro Gln Asp Ala Ser

1502 1511 1520 1529 1538 GTA TCT CCT GCA ATT GGG TAT AGT CAC CGT TTA TGC CAT Val Ser Pro Ala Ile Gly Tyr Ser His Arg Leu Cys His

1547 1556 1565 1574

GCA ACA TTT TTA GAA CGG ATT AGT GGA CCA AGA ATA GCA
Ala Thr Phe Leu Glu Arg Ile Ser Gly Pro Arg Ile Ala

1583 1592 1601 1610

GGC ACC GTA TTT TCT TGG ACA CAC CGT AGT GCC AGC CCT

Gly Thr Val Phe Ser Trp Thr His Arg Ser Ala Ser Pro

1619 1628 1637 1646 1655 ACT AAT GAA GTA AGT CCA TCT AGA ATT ACA CAA ATT CCA Thr Asn Glu Val Ser Pro Ser Arg Ile Thr Gln Ile Pro

1664 1673 1682 1691
TGG GTA AAG GCG CAT ACT CTT GCA TCT GGT GCC TCC GTC
Trp Val Lys Ala His Thr Leu Ala Ser Gly Ala Ser Val

Figure 13 (Cont. 7)

1700 1709 1718 1727
ATT AAA GGT CCT GGA TTT ACA GGT GGA GAT ATT CTG ACT
Ile Lys Gly Pro Gly Phe Thr Gly Gly Asp Ile Leu Thr

1736 1745 1754 1763 1772

AGG AAT AGT ATG GGC GAG CTG GGG ACC TTA CGA GTA ACC

Arg Asn Ser MET Gly Glu Leu Gly Thr Leu Arg Val Thr

1781 1790 1799 1808
TTC ACA GGA AGA TTA CCA CAA AGT TAT TAT ATA CGT TTC
Phe Thr Gly Arg Leu Pro Gln Ser Tyr Tyr Ile Arg Phe

1817 1826 1835 1844

CGT TAT GCT TCG GTA GCA AAT AGG AGT GGT ACA TTT AGA

Arg Tyr Ala Ser Val Ala Asn Arg Ser Gly Thr Phe Arg

1853 1862 1871 1880 1889
TAT TCA CAG CCA CCT TCG TAT GGA ATT TCA TTT CCA AAA
Tyr Ser Gln Pro Pro Ser Tyr Gly Ile Ser Phe Pro Lys

1898 1907 1916 1925
ACT ATG GAC GCA GGT GAA CCA CTA ACA TCT CGT TCG TTC
Thr MET Asp Ala Gly Glu Pro Leu Thr Ser Arg Ser Phe

Figure 13 (Cont. 8)

GCT CAT ACA ACA CTC TTC ACT CCA ATA ACC TTT TCA CGA Ala His Thr Thr Leu Phe Thr Pro Ile Thr Phe Ser Arg GCT CAA GAA GAA TTT GAT CTA TAC ATC CAA TCG GGT GTT Ala Gln Glu Glu Phe Asp Leu Tyr Ile Gln Ser Gly Val TAT ATA GAT CGA ATT GAA TTT ATA CCG GTT ACT GCA ACA Tyr Ile Asp Arg Ile Glu Phe Ile Pro Val Thr Ala Thr TTT GAG GCA GAA TAT GAT TTA GAA AGA GCG CAA AAG GTG Phe Glu Ala Glu Tyr Asp Leu Glu Arg Ala Gln Lys Val GTG AAT GCC CTG TTT ACG TCT ACA AAC CAA CTA GGG CTA Val Asn Ala Leu Phe Thr Ser Thr Asn Gln Leu Gly Leu AAA ACA GAT GTG ACG GAT TAT CAT ATT GAT CAG GTA TCC Lys Thr Asp Val Thr Asp Tyr His Ile Asp Gln Val Ser

Figure 13 (Cont. 9)

2168 2177 2186 2195

AAT CTA GTT GCG TGT TTA TCG GAT GAA TTT TGT CTG GAT

Asn Leu Val Ala Cys Leu Ser Asp Glu Phe Cys Leu Asp

2204 2213 2222 2231 2240 GAA AAG AGA GAA TTG TCC GAG AAA GTT AAA CAT GCA AAG Glu Lys Arg Glu Leu Ser Glu Lys Val Lys His Ala Lys

2249 2258 2267 2276
CGA CTC AGT GAT GAG CGG AAT TTA CTT CAA GAT CCA AAC
Arg Leu Ser Asp Glu Arg Asn Leu Leu Gln Asp Pro Asn

2285 2294 2303 2312

TTC AGA GGG ATC AAT AGG CAA CCA GAC CGT GGC TGG AGA

Phe Arg Gly Ile Asn Arg Gln Pro Asp Arg Gly Trp Arg

2321 2330 2339 2348 2357 GGA AGT ACG GAT ATT ACT ATC CAA GGA GGA GAT GAC GTA Gly Ser Thr Asp Ile Thr Ile Gln Gly Gly Asp Asp Val

2366 2375 2384 2393
TTC AAA GAG AAT TAC GTT ACG CTA CCG GGT ACC TTT GAT
Phe Lys Glu Asn Tyr Val Thr Leu Pro Gly Thr Phe Asp

Figure 13 (Cont. 10)

2402 2411 2420 2429

GAG TGC TAT CCA ACG TAT TTA TAT CAA AAA ATA GAT GAG
Glu Cys Tyr Pro Thr Tyr Leu Tyr Gln Lys Ile Asp Glu

2438 2447 2456 2465 2474
TCG AAA TTA AAA GCC TAT ACC CGT TAT CAA TTA AGA GGG
Ser Lys Leu Lys Ala Tyr Thr Arg Tyr Gln Leu Arg Gly

2483 2492 2501 2510
TAT ATC GAA GAT AGT CAA GAC TTA GAA ATC TAT TTA ATT
Tyr Ile Glu Asp Ser Gln Asp Leu Glu Ile Tyr Leu Ile

2519 2528 2537 2546

CGT TAC AAT GCA AAA CAC GAA ATA GTA AAT GTA CCA GGT

Arg Tyr Asn Ala Lys His Glu Ile Val Asn Val Pro Gly

2555 2564 2573 2582 2591
ACA GGA AGT TTA TGG CCT CTT TCT GTA GAA AAT CAA ATT
Thr Gly Ser Leu Trp Pro Leu Ser Val Glu Asn Gln Ile

2600 2609 2618 2627

GGA CCT TGT GGA GAA CCG AAT CGA TGC GCG CCA CAC CTT

Gly Pro Cys Gly Glu Pro Asn Arg Cys Ala Pro His Leu

Figure 13 (Cont. 11)

2636 2645 2654 2663

GAA TGG AAT CCT GAT TTA CAC TGT TCC TGC AGA GAC GGG
Glu Trp Asn Pro Asp Leu His Cys Ser Cys Arg Asp Gly

2672 2681 2690 2699 2708
GAA AAA TGT GCA CAT CAT TCT CAT CAT TTC TCT TTG GAC
Glu Lys Cys Ala His His Ser His His Phe Ser Leu Asp

2717 2726 2735 2744

ATT GAT GTT GGA TGT ACA GAC TTA AAT GAG GAC TTA GGT

Ile Asp Val Gly Cys Thr Asp Leu Asn Glu Asp Leu Gly

2753 2762 2771 2780

GTA TGG GTG ATA TTC AAG ATT AAG ACG CAA GAT GGC CAC

Val Trp Val Ile Phe Lys Ile Lys Thr Gln Asp Gly His

2789 2798 2807 . 2816 2825 GCA CGA CTA GGG AAT CTA GAG TTT CTC GAA GAG AAA CCA Ala Arg Leu Gly Asn Leu Glu Phe Leu Glu Glu Lys Pro

2834 2843 2852 2861
TTA TTA GGA GAA GCA CTA GCT CGT GTG AAA AGA GCG GAG
Leu Leu Gly Glu Ala Leu Ala Arg Val Lys Arg Ala Glu

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Figure 13 (Cont. 12)

2870 2879 2888 2897

AAA AAA TGG AGA GAC AAA CGC GAA ACA TTA CAA TTG GAA
Lys Lys Trp Arg Asp Lys Arg Glu Thr Leu Gln Leu Glu

2906 2915 2924 2933 2942 ACA ACT ATC GTT TAT AAA GAG GCA AAA GAA TCT GTA GAT Thr Thr Ile Val Tyr Lys Glu Ala Lys Glu Ser Val Asp

2951 2960 2969 2978
GCT TTA TTT GTA AAC TCT CAA TAT GAT AGA TTA CAA GCG
Ala Leu Phe Val Asn Ser Gln Tyr Asp Arg Leu Gln Ala

2987 2996 3005 3014

GAT ACG AAC ATC GCG ATG ATT CAT GCG GCA GAT AAA CGC
Asp Thr Asn Ile Ala MET Ile His Ala Ala Asp Lys Arg

3023 3032 3041 3050 3059 GTT CAT AGA ATT CGA GAA GCG TAT CTG CCG GAG CTG TCT Val His Arg Ile Arg Glu Ala Tyr Leu Pro Glu Leu Ser

3068 3077 3086 3095
GTG ATT CCG GGT GTC AAT GCG GCT ATT TTT GAA GAA TTA
Val Ile Pro Gly Val Asn Ala Ala Ile Phe Glu Glu Leu

Figure 13 (Cont. 13)

3104 3113 3122 3131

GAA GAG CGT ATT TTC ACT GCA TTT TCC CTA TAT GAT GCG
Glu Glu Arg Ile Phe Thr Ala Phe Ser Leu Tyr Asp Ala

3140 3149 3158 3167 3176

AGA AAT ATT ATT AAA AAT GGC GAT TTC AAT AAT GGC TTA

Arg Asn Ile Ile Lys Asn Gly Asp Phe Asn Asn Gly Leu

3185 3194 3203 3212
TTA TGC TGG AAC GTG AAA GGG CAT GTA GAG GTA GAA GAA
Leu Cys Trp Asn Val Lys Gly His Val Glu Val Glu Glu

3221 3230 3239 3248

CAA AAC AAT CAC CGT TCA GTC CTG GTT ATC CCA GAA TGG
Gln Asn Asn His Arg Ser Val Leu Val Ile Pro Glu Trp

3257 3266 3275 3284 3293 GAG GCA GAA GTG TCA CAA GAG GTT CGT GTC TGT CCA GGT Glu Ala Glu Val Ser Gln Glu Val Arg Val Cys Pro Gly

3302 3311 3320 3329

CGT GGC TAT ATC CTT CGT GTT ACA GCG TAC AAA GAG GGA

Arg Gly Tyr Ile Leu Arg Val Thr Ala Tyr Lys Glu Gly

Figure 13 (Cont. 14)

3338 3347 3356 3365

TAT GGA GAA GGT TGC GTA ACG ATC CAT GAG ATC GAG AAC

Tyr Gly Glu Gly Cys Val Thr Ile His Glu Ile Glu Asn

3374 3383 3392 3401 3410
AAT ACA GAC GAA CTG AAA TTC AAC AAC TGT GTA GAA GAG
Asn Thr Asp Glu Leu Lys Phe Asn Asn Cys Val Glu Glu

3419 3428 3437 3446

GAA GTA TAT CCA AAC AAC ACG GTA ACG TGT ATT AAT TAT
Glu Val Tyr Pro Asn Asn Thr Val Thr Cys Ile Asn Tyr

3455 3464 3473 3482

ACT GCG ACT CAA GAA GAA TAT GAG GGT ACG TAC ACT TCT

Thr Ala Thr Gln Glu Glu Tyr Glu Gly Thr Tyr Thr Ser

3491 3500 3509 3518 3527 CGT AAT CGA GGA TAT GAC GAA GCC TAT GGT AAT AAC CCT Arg Asn Arg Gly Tyr Asp Glu Ala Tyr Gly Asn Asn Pro

3536 3545 3554 3563
TCC GTA CCA GCT GAT TAT GCG TCA GTC TAT GAA GAA AAA
Ser Val Pro Ala Asp Tyr Ala Ser Val Tyr Glu Glu Lys

Figure 13 (Cont. 15)

3572 3581 3590 3599
TCG TAT ACA GAT AGA CGA AGA GAG AAT CCT TGT GAA TCT
Ser Tyr Thr Asp Arg Arg Glu Asn Pro Cys Glu Ser

3608 3617 3626 3635 3644
AAC AGA GGA TAT GGA GAT TAC ACA CCA CTA CCA GCT GGT
Asn Arg Gly Tyr Gly Asp Tyr Thr Pro Leu Pro Ala Gly

3653 3662 3671 3680

TAT GTA ACA AAG GAA TTA GAG TAC TTC CCA GAG ACC GAT

Tyr Val Thr Lys Glu Leu Glu Tyr Phe Pro Glu Thr Asp

3689 3698 3707 3716

AAG GTA TGG ATT GAG ATT GGA GAA ACA GAA GGA ACA TTC
Lys Val Trp Ile Glu Ile Gly Glu Thr Glu Gly Thr Phe

3725 3734 3743 3752 3761
ATC GTG GAC AGC GTG GAA TTA CTC CTT ATG GAG GAA TAG
Ile Val Asp Ser Val Glu Leu Leu Leu MET Glu Glu

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Figure 13 (Cont. 16)

3771 3781 3791 3801 3811 GACCATCCGA GTATAGCAGT TTAATAAATA TTAATTAAAA TAGTAGTCTA

3821 3831 3841 3851 3861 ACTTCCGTTC CAATTAAATA AGTAAATTAC AGTTGTAAAA AAAAACGAAC

3871 3881 3891 3901 ATTACTCTTC AAAGAGCGAT GTCCGTTTTT TATATGGTGT GT

Figure 14

10	20	30	40	
AATAGAATCT	CAAATCTCGA	TGACTGCTTA	GTCTTTTTAA	
60 TTGACAGGGG		80 ATCGGTCAAT		100 GGGCATATAT
110	120	130	140	150
TGATATTTTA	TAAAATTTGT	TACGTTTTTT	GTATTTTTC	ATAAGATGTG
160	- 1 0	180	190	200
TCATATGTAT		TAATGAAAAA	CAGTATCAAA	CTATCAGAAC
	220 TAATAAAAA	230 ACGGAGGTAT 		GAA
248 AAT AAT CAA Asn Asn Gln	AAT CAA TG	266 C ATA CCT T s Ile Pro T	AC AAT TGT	TTA AGT Leu Ser
284	GAA GTA CT	302	311	320
AAT CCT GAA		T TTG GAT GO	GA GAA CGG A	ATA TCA
Asn Pro Glu		u Leu Asp G	ly Glu Arg 1	Ile Ser

Figure 14 (Cont. 1)

		AAT	TCA	TCA	ATT	GAT	ATT	TCT	CTG	TCA	CTT	
Thr	Gly	Asn	Ser	Ser	Ile	Asp	Ile	Ser	Leu	Ser	Leu	Val
	365			374			383			392		
				TCT								
Gln	Phe	Leu	Val	Ser	Asn	Phe	Val	Pro	Gly	Gly	Gly	Phe
401			410			419			428			437
TTA	GTT	GGA	TTA	ATA	GAT	TTT	GTA	TGG	GGA	ATA	GTT	GGC
Leu	Val	Gly	Leu	Ile	Asp	Phe	Val	Trp	Gly	Ile	Val	Gly
		446			455			464			473	
CCT	TCT			GAT								CAA
		CAA	TGG	GAT	GCA	TTT	CTA	GTA	CAA	ATT	GAA	
	Ser	CAA Gln	TGG Trp	GAT Asp	GCA Ala	TTT Phe	CTA Leu	GTA Val	CAA Gln	ATT Ile	GAA	
Pro	Ser 482	CAA Gln	TGG Trp	GAT Asp 491	GCA Ala	TTT Phe	CTA Leu 500	GTA Val	CAA Gln	ATT Ile 509	GAA Glu	Gln
Pro TTA	Ser 482 ATT	CAA Gln AAT	TGG Trp GAA	GAT Asp 491 AGA	GCA Ala ATA	TTT Phe GCT	CTA Leu 500 GAA	GTA Val	CAA Gln GCT	ATT Ile 509 AGG	GAA Glu AAT	Gln GCT
Pro TTA	Ser 482 ATT	CAA Gln AAT	TGG Trp GAA	GAT Asp 491	GCA Ala ATA	TTT Phe GCT	CTA Leu 500 GAA	GTA Val	CAA Gln GCT	ATT Ile 509 AGG	GAA Glu AAT	Gln GCT
Pro TTA	Ser 482 ATT	CAA Gln AAT	TGG Trp GAA	GAT Asp 491 AGA Arg	GCA Ala ATA	TTT Phe GCT	CTA Leu 500 GAA Glu	GTA Val	CAA Gln GCT	ATT Ile 509 AGG	GAA Glu AAT	Gln GCT
Pro TTA Leu 518 GCT	Ser 482 ATT Ile	CAA Gln AAT Asn GCT	TGG Trp GAA Glu 527 AAT	GAT Asp 491 AGA Arg	GCA Ala ATA Ile GAA	TTT Phe GCT Ala 536 GGA	CTA Leu 500 GAA Glu TTA	GTA Val TTT Phe	CAA Gln GCT Ala 545 AAC	ATT Ile 509 AGG Arg	GAA Glu AAT Asn	Gln GCT Ala 554 AAT

Figure 14 (Cont. 2)

		563			572			581			590	
ATA	TAT	GTG	GAA	GCA	TTT	AAA	GAA	TGG	GAA	GAA	GAT	CCT
				Ala								
	599			608			617			626	•	
AAT				ACC							CGC-	TTT
				Thr								
635			644			653			662	·		671
CGT	ATA	CTT	GAT	GGG	CTA	CTT	GAA	AGG	GAC	ATT	CCT	TCG
				Gly								
		680			689			698			707	
TTT	CGA			GGA								
Phe	Arg	Ile	Ser	Gly	Phe	Glu	Val	Pro	Leu	Leu	Ser	Val
	716			725			734			743		
TAT				GCC							TTA	AGA
Tyr	Ala	Gln	Ala	Ala	Asn	Leu	His	Leu	Ala	Ile	Leu	Arg
752			761			770			779			788
GAT	TCT	GTA	ATT	TTT	GGA	GAA	AGA	TGG	GGA	TTG	ACA	ACG
Asp	Ser	Val	Ile	Phe	Gly	Glu	Arg	Trp	Gly	Leu	Thr	Thr

Figure 14 (Cont. 3) 806 815 824 797 ATA AAT GTC AAT GAA AAC TAT AAT AGA CTA ATT AGG CAT Ile Asn Val Asn Glu Asn Tyr Asn Arg Leu Ile Arg His 842 833 851 860 -ATT GAT GAA TAT GCT GAT CAC TGT GCA AAT ACG TAT AAT Ile Asp Glu Tyr Ala Asp His Cys Ala Asn Thr Tyr Asn 869 878 887 896 905 CGG GGA TTA AAT AAT TTA CCG AAA TCT ACG TAT CAA GAT Arg Gly Leu Asn Asn Leu Pro Lys Ser Thr Tyr Gln Asp 914 923 932 941 TGG ATA ACA TAT AAT CGA TTA CGG AGA GAC TTA ACA TTG Trp Ile Thr Tyr Asn Arg Leu Arg Arg Asp Leu Thr Leu 950 959 968 977 ACT GTA TTA GAT ATC GCC GCT TTC TTT CCA AAC TAT GAC Thr Val Leu Asp Ile Ala Ala Phe Phe Pro Asn Tyr Asp 986 995 1004 1013 1022 AAT AGG AGA TAT CCA ATT CAG CCA GTT GGT CAA CTA ACA Asn Arg Arg Tyr Pro Ile Gln Pro Val Gly Gln Leu Thr

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Figure 14 (Cont. 4)

1031 1040 1049 1058
AGG GAA GTT TAT ACG GAC CCA TTA ATT AAT TTT AAT CCA
Arg Glu Val Tyr Thr Asp Pro Leu Ile Asn Phe Asn Pro

1067 1076 1085 1094
CAG TTA CAG TCT GTA GCT CAA TTA CCT ACT TTT AAC GTT
Gln Leu Gln Ser Val Ala Gln Leu Pro Thr Phe Asn Val

1103 1112 1121 1130 1139
ATG GAG AGC AGC GCA ATT AGA AAT CCT CAT TTA TTT GAT
MET Glu Ser Ser Ala Ile Arg Asn Pro His Leu Phe Asp

1148 1157 1166 1175
ATA TTG AAT AAT CTT ACA ATC TTT ACG GAT TGG TTT AGT
Ile Leu Asn Asn Leu Thr Ile Phe Thr Asp Trp Phe Ser

1184 1193 1202 1211
GTT GGA CGC AAT TTT TAT TGG GGA GGA CAT CGA GTA ATA
Val Gly Arg Asn Phe Tyr Trp Gly Gly His Arg Val Ile

1220 1229 1238 1247 1256

TCT AGC CTT ATA GGA GGT GGT AAC ATA ACA TCT CCT ATA Ser Ser Leu Ile Gly Gly Gly Asn Ile Thr Ser Pro Ile

Figure 14 (Cont. 5)

1265 1274 1283 1292
TAT GGA AGA GAG GCG AAC CAG GAG CCT CCA AGA TCC TTT
Tyr Gly Arg Glu Ala Asn Gln Glu Pro Pro Arg Ser Phe

1301 1310 1319 1328
ACT TTT AAT GGA CCG GTA TTT AGG ACT TTA TCA AAT CCT
Thr Phe Asn Gly Pro Val Phe Arg Thr Leu Ser Asn Pro

1337 1346 1355 1364 1373
ACT TTA CGA TTA TTA CAG CAA CCT TGG CCA GCG CCA CCA
Thr Leu Arg Leu Leu Gln Gln Pro Trp Pro Ala Pro Pro

1382 1391 1400 1409
TTT AAT TTA CGT GGT GTT GAA GGA GTA GAA TTT TCT ACA
Phe Asn Leu Arg Gly Val Glu Gly Val Glu Phe Ser Thr

1418 1427 1436 1445
CCT ACA AAT AGC TTT ACG TAT CGA GGA AGA GGT ACG GTT
Pro Thr Asn Ser Phe Thr Tyr Arg Gly Arg Gly Thr Val

1454 1463 1472 1481 1490 GAT TCT TTA ACT GAA TTA CCG CCT GAG GAT AAT AGT GTG Asp Ser Leu Thr Glu Leu Pro Pro Glu Asp Asn Ser Val

Figure 14 (Cont. 6)

1499 1508 1517 1526
CCA CCT CGC GAA GGA TAT AGT CAT CGT TTA TGT CAT GCA
Pro Pro Arg Glu Gly Tyr Ser His Arg Leu Cys His Ala

1535 1544 1553 1562
ACT TTT GTT CAA AGA TCT GGA ACA CCT TTT TTA ACA ACT
Thr Phe Val Gln Arg Ser Gly Thr Pro Phe Leu Thr Thr

1571 1580 1589 1598 1607 GGT GTA GTA TTT TCT TGG ACG CAT CGT AGT GCA ACT CTT Gly Val Val Phe Ser Trp Thr His Arg Ser Ala Thr Leu

1616 1625 1634 1643
ACA AAT ACA ATT GAT CCA GAG AGA ATT AAT CAA ATA CCT
Thr Asn Thr Ile Asp Pro Glu Arg Ile Asn Gln Ile Pro

1652 1661 1670 1679
TTA GTG AAA GGA TTT AGA GTT TGG GGG GGC ACC TCT GTC
Leu Val Lys Gly Phe Arg Val Trp Gly Gly Thr Ser Val

1688 1697 1706 1715 1724
ATT ACA GGA CCA GGA TTT ACA GGA GGG GAT ATC CTT CGA
Ile Thr Gly Pro Gly Phe Thr Gly Gly Asp Ile Leu Arg

Figure 14 (Cont. 7)

1733 1742 1751 1760
AGA AAT ACC TTT GGT GAT TTT GTA TCT CTA CAA GTC AAT
Arg Asn Thr Phe Gly Asp Phe Val Ser Leu Gln Val Asn

1769 1778 1787 1796
ATT AAT TCA CCA ATT ACC CAA AGA TAC CGT TTA AGA TTT
Ile Asn Ser Pro Ile Thr Gln Arg Tyr Arg Leu Arg Phe

1805 1814 1823 1832 1841 CGT TAC GCT TCC AGT AGG GAT GCA CGA GTT ATA GTA TTA Arg Tyr Ala Ser Ser Arg Asp Ala Arg Val Ile Val Leu

1850 1859 1868 1877
ACA GGA GCG GCA TCC ACA GGA GTG GGA GGC CAA GTT AGT
Thr Gly Ala Ala Ser Thr Gly Val Gly Gly Gln Val Ser

1886 1895 1904 1913
GTA AAT ATG CCT CTT CAG AAA ACT ATG GAA ATA GGG GAG
Val Asn MET Pro Leu Gln Lys Thr MET Glu Ile Gly Glu

1922 1931 1940 1949 1958
AAC TTA ACA TCT AGA ACA TTT AGA TAT ACC GAT TTT AGT
Asn Leu Thr Ser Arg Thr Phe Arg Tyr Thr Asp Phe Ser

Figure 14 (Cont. 8)

1967 1976 1985 1994

AAT CCT TTT TCA TTT AGA GCT AAT CCA GAT ATA ATT GGG

Asn Pro Phe Ser Phe Arg Ala Asn Pro Asp Ile Ile Gly

2003 2012 2021 2030
ATA AGT GAA CAA CCT CTA TTT GGT GCA GGT TCT ATT AGT
Ile Ser Glu Gln Pro Leu Phe Gly Ala Gly Ser Ile Ser

2039 2048 2057 2066 2075
AGC GGT GAA CTT TAT ATA GAT AAA ATT GAA ATT ATT CTA
Ser Gly Glu Leu Tyr Ile Asp Lys Ile Glu Ile Ile Leu

2084 2093 2102 2111
GCA GAT GCA ACA TTT GAA GCA GAA TCT GAT TTA GAA AGA
Ala Asp Ala Thr Phe Glu Ala Glu Ser Asp Leu Glu Arg

2120 2129 2138 2147
GCA CAA AAG GCG GTG AAT GCC CTG TTT ACT TCT TCC AAT
Ala Gln Lys Ala Val Asn Ala Leu Phe Thr Ser Ser Asn

2156 2165 2174 2183 2192 CAA ATC GGG TTA AAA ACC GAT GTG ACG GAT TAT CAT ATT Gln Ile Gly Leu Lys Thr Asp Val Thr Asp Tyr His Ile

Figure 14 (Cont. 9)

2201 2210 2219 2228

GAT CAA GTA TCC AAT TTA GTG GAT TGT TTA TCA GAT GAA

Asp Gln Val Ser Asn Leu Val Asp Cys Leu Ser Asp Glu

2237 2246 2255 2264
TTT TGT CTG GAT GAA AAG CGA GAA TTG TCC GAG AAA GTC
Phe Cys Leu Asp Glu Lys Arg Glu Leu Ser Glu Lys Val

2273 2282 2291 2300 2309
AAA CAT GCG AAG CGA CTC AGT GAT GAG CGG AAT TTA CTT
Lys His Ala Lys Arg Leu Ser Asp Glu Arg Asn Leu Leu

2318 2327 2336 2345
CAA GAT CCA AAC TTC AGA GGG ATC AAT AGA CAA CCA GAC
Gln Asp Pro Asn Phe Arg Gly Ile Asn Arg Gln Pro Asp

2354 2363 2372 2381
CGT GGC TGG AGA GGA AGT ACA GAT ATT ACC ATC CAA GGA
Arg Gly Trp Arg Gly Ser Thr Asp Ile Thr Ile Gln Gly

2390 2399 2408 2417 2426 GGA GAT GAC GTA TTC AAA GAG AAT TAC GTC ACA CTA CCG Gly Asp Asp Val Phe Lys Glu Asn Tyr Val Thr Leu Pro

Figure 14 (Cont. 10)

2435 2444 2453 2462

GGT ACC GTT GAT GAG TGC TAT CCA ACG TAT TTA TAT CAG

Gly Thr Val Asp Glu Cys Tyr Pro Thr Tyr Leu Tyr Gln

2471 2480 2489 2498

AAA ATA GAT GAG TCG AAA TTA AAA GCT TAT ACC CGT TAT
Lys Ile Asp Glu Ser Lys Leu Lys Ala Tyr Thr Arg Tyr

2507 2516 2525 2534 2543 GAA TTA AGA GGG TAT ATC GAA GAT AGT CAA GAC TTA GAA Glu Leu Arg Gly Tyr Ile Glu Asp Ser Gln Asp Leu Glu

2552 2561 2570 2579

ATC TAT TTG ATC CGT TAC AAT GCA AAA CAC GAA ATA GTA

Ile Tyr Leu Ile Arg Tyr Asn Ala Lys His Glu Ile Val

2588 2597 2606 2615

AAT GTG CCA GGC ACG GGT TCC TTA TGG CCG CTT TCA GCC

Asn Val Pro Gly Thr Gly Ser Leu Trp Pro Leu Ser Ala

2624 2633 2642 2651 2660 CAA AGT CCA ATC GGA AAG TGT GGA GAA CCG AAT CGA TGC Gln Ser Pro Ile Gly Lys Cys Gly Glu Pro Asn Arg Cys

Figure 14 (Cont. 11)

2669 2678 2687 2696

GCG CCA CAC CTT GAA TGG AAT CCT GAT CTA GAT TGT TCC

Ala Pro His Leu Glu Trp Asn Pro Asp Leu Asp Cys Ser

2705 2714 2723 2732
TGC AGA GAC GGG GAA AAA TGT GCA CAT CAT TCC CAT CAT
Cys Arg Asp Gly Glu Lys Cys Ala His His Ser His His

2741 2750 2759 2768 2777
TTC ACC TTG GAT ATT GAT GTT GGA TGT ACA GAC TTA AAT
Phe Thr Leu Asp Ile Asp Val Gly Cys Thr Asp Leu Asn

2786 2795 2804 2813

GAG GAC TTA GGT GTA TGG GTG ATA TTC AAG ATT AAG ACG
Glu Asp Leu Gly Val Trp Val Ile Phe Lys Ile Lys Thr

2822 2831 2840 2849
CAA GAT GGC CAT GCA AGA CTA GGG AAT CTA GAG TTT CTC
Gln Asp Gly His Ala Arg Leu Gly Asn Leu Glu Phe Leu

2858 2867 2876 2885 2894
GAA GAG AAA CCA TTA TTA GGG GAA GCA CTA GCT CGT GTG
Glu Glu Lys Pro Leu Leu Gly Glu Ala Leu Ala Arg Val

Figure 14 (Cont. 12)

2903 2912 2921 2930
AAA AGA GCG GAG AAG AAG TGG AGA GAC AAA CGA GAG AAA
Lys Arg Ala Glu Lys Lys Trp Arg Asp Lys Arg Glu Lys

2939 2948 2957 2966
CTG CAG TTG GAA ACA AAT ATT GTT TAT AAA GAG GCA AAA
Leu Gln Leu Glu Thr Asn Ile Val Tyr Lys Glu Ala Lys

2975 2984 2993 3002 3011 GAA TCT GTA GAT GCT TTA TTT GTA AAC TCT CAA TAT GAT Glu Ser Val Asp Ala Leu Phe Val Asn Ser Gln Tyr Asp

3020 3029 3038 3047
AGA TTA CAA GTG GAT ACG AAC ATC GCG ATG ATT CAT GCG
Arg Leu Gln Val Asp Thr Asn Ile Ala MET Ile His Ala

3056 3065 3074 3083

GCA GAT AAA CGC GTT CAT AGA ATC CGG GAA GCG TAT CTG

Ala Asp Lys Arg Val His Arg Ile Arg Glu Ala Tyr Leu

3092 3101 3110 3119 3128 CCA GAG TTG TCT GTG ATT CCA GGT GTC AAT GCG GCC ATT Pro Glu Leu Ser Val Ile Pro Gly Val Asn Ala Ala Ile

Figure 14 (Cont. 13)

3137 3146 3155 3164
TTC GAA GAA TTA GAG GGA CGT ATT TTT ACA GCG TAT TCC
Phe Glu Glu Leu Glu Gly Arg Ile Phe Thr Ala Tyr Ser

3173 3182 3191 3200
TTA TAT GAT GCG AGA AAT GTC ATT AAA AAT GGC GAT TTC
Leu Tyr Asp Ala Arg Asn Val Ile Lys Asn Gly Asp Phe

3209 3218 3227 3236 3245
AAT AAT GGC TTA TTA TGC TGG AAC GTG AAA GGT CAT GTA
Asn Asn Gly Leu Leu Cys Trp Asn Val Lys Gly His Val

3254 3263 3272 3281
GAT GTA GAA GAG CAA AAC AAC CAC CGT TCG GTC CTT GTT
Asp Val Glu Glu Gln Asn Asn His Arg Ser Val Leu Val

3290 3299 3308 3317
ATC CCA GAA TGG GAG GCA GAA GTG TCA CAA GAG GTT CGT
Ile Pro Glu Trp Glu Ala Glu Val Ser Gln Glu Val Arg

3326 3335 3344 3353 3362
GTC TGT CCA GGT CGT GGC TAT ATC CTT CGT GTC ACA GCA
Val Cys Pro Gly Arg Gly Tyr Ile Leu Arg Val Thr Ala

Figure 14 (Cont. 14)

3371 3380 3389 3398
TAT AAA GAG GGA TAT GGA GAG GGC TGC GTA ACG ATC CAT
Tyr Lys Glu Gly Tyr Gly Glu Gly Cys Val Thr Ile His

3407 3416 3425 3434

GAG ATC GAA GAC AAT ACA GAC GAA CTG AAA TTC AGC AAC
Glu Ile Glu Asp Asn Thr Asp Glu Leu Lys Phe Ser Asn

3443 3452 3461 3470 3479
TGT GTA GAA GAG GAA GTA TAT CCA AAC AAC ACA GTA ACG
Cys Val Glu Glu Val Tyr Pro Asn Asn Thr Val Thr

3488 3497 3506 3515
TGT AAT AAT TAT ACT GGG ACT CAA GAA GAA TAT GAG GGT
Cys Asn Asn Tyr Thr Gly Thr Gln Glu Glu Tyr Glu Gly

3524 3533 3542 3551

ACG TAC ACT TCT CGT AAT CAA GGA TAT GAC GAA GCC TAT

Thr Tyr Thr Ser Arg Asn Gln Gly Tyr Asp Glu Ala Tyr

3560 3569 3578 3587 3596
GGT AAT AAC CCT TCC GTA CCA GCT GAT TAC GCT TCA GTC
Gly Asn Asn Pro Ser Val Pro Ala Asp Tyr Ala Ser Val

Figure 14 (Cont. 15)

3605 3614 3623 3632

TAT GAA GAA AAA TCG TAT ACA GAT GGA CGA AGA GAG AAT

Tyr Glu Glu Lys Ser Tyr Thr Asp Gly Arg Arg Glu Asn

3641 3650 3659 3668
CCT TGT GAA TCT AAC AGA GGC TAT GGG GAT TAC ACA CCA
Pro Cys Glu Ser Asn Arg Gly Tyr Gly Asp Tyr Thr Pro

3677 3686 3695 3704 3713 CTA CCG GCT GGT TAT GTA ACA AAG GAT TTA GAG TAC TTC Leu Pro Ala Gly Tyr Val Thr Lys Asp Leu Glu Tyr Phe

3722 3731 3740 3749
CCA GAG ACC GAT AAG GTA TGG ATT GAG ATC GGA GAA ACA
Pro Glu Thr Asp Lys Val Trp Ile Glu Ile Gly Glu Thr

3758 3767 3776 3785

GAA GGA ACA TTC ATC GTG GAT AGC GTG GAA TTA CTC CTT

Glu Gly Thr Phe Ile Val Asp Ser Val Glu Leu Leu Leu

3794 3803 3813 3823 3833 ATG GAG GAA TAA GATACGTTAT AAAATGTAAC GTATGCAAAT MET Glu Glu .

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Figure 14 (Cont. 16)

3843 3853 3863 3873 3883 AAAGAATGAT TACTGACCTA TATTAACAGA TAAATAAGAA AATTTTTATA

3893 3903 3913 3923 CGAATAAAAA ACGGACATCA CTCTTAAGAG AATGATGTCC

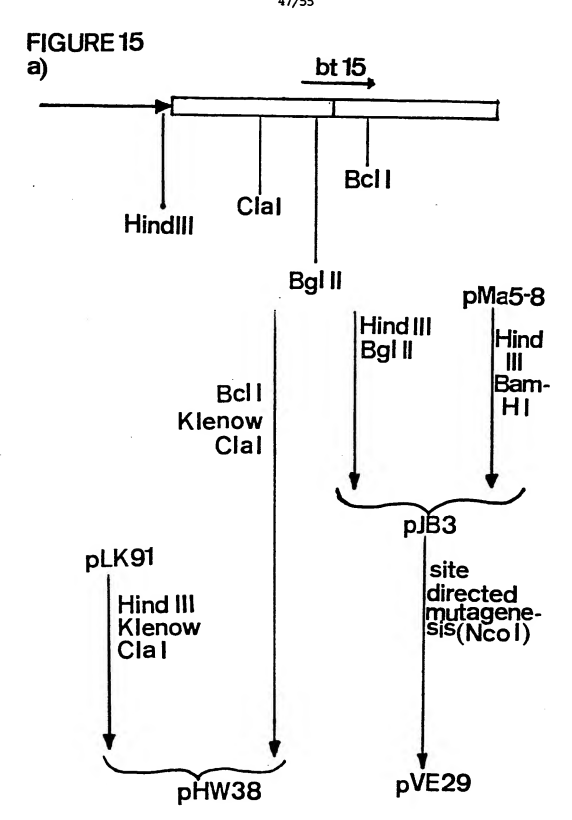
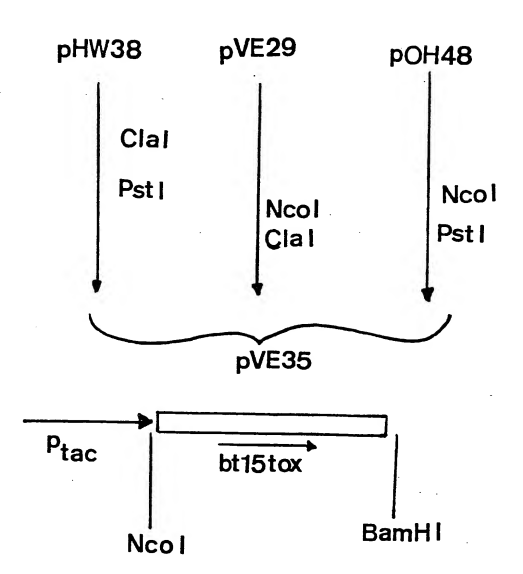


Fig. 15 (cont)

b)



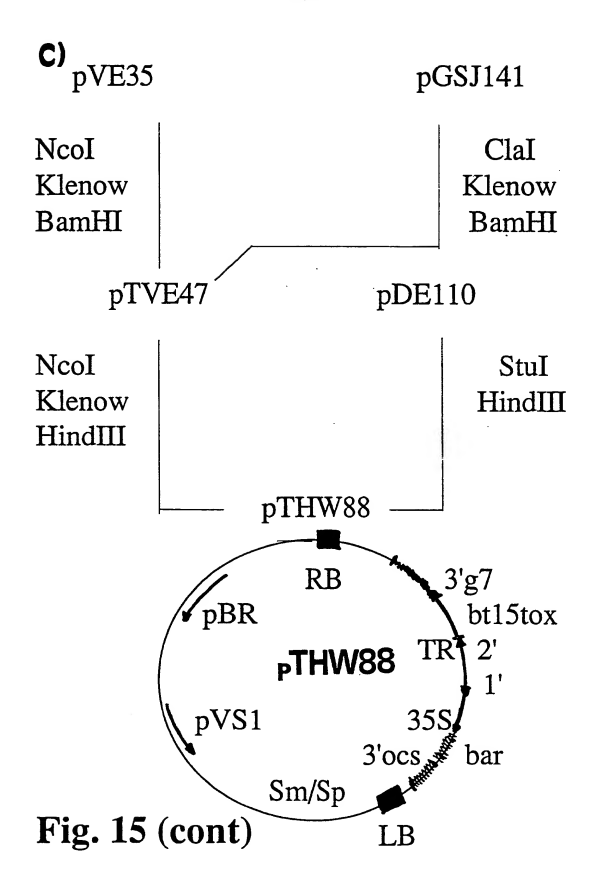
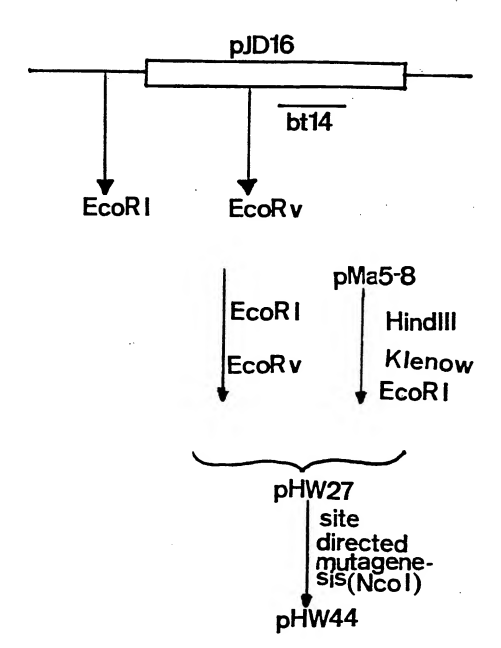
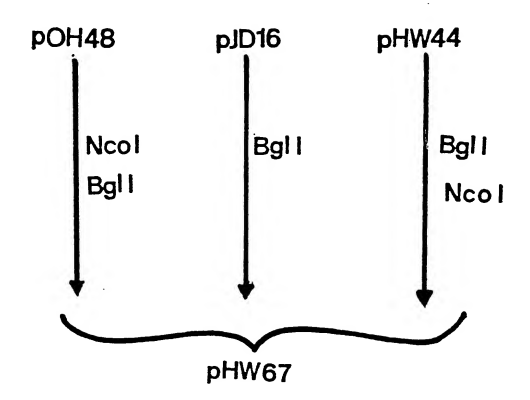


FIGURE 16 a)



### FIGURE 16 b)

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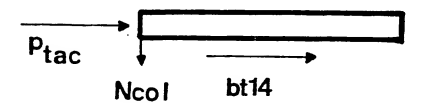


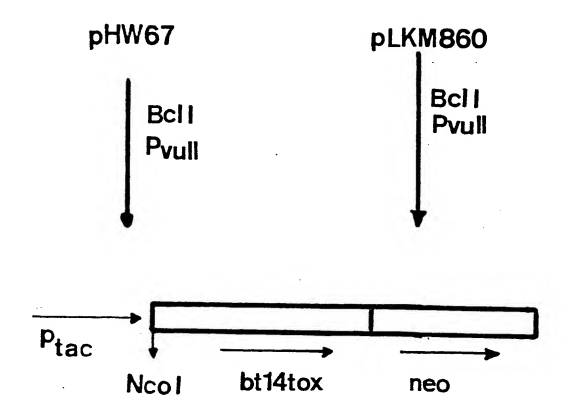
FIGURE 16

c)

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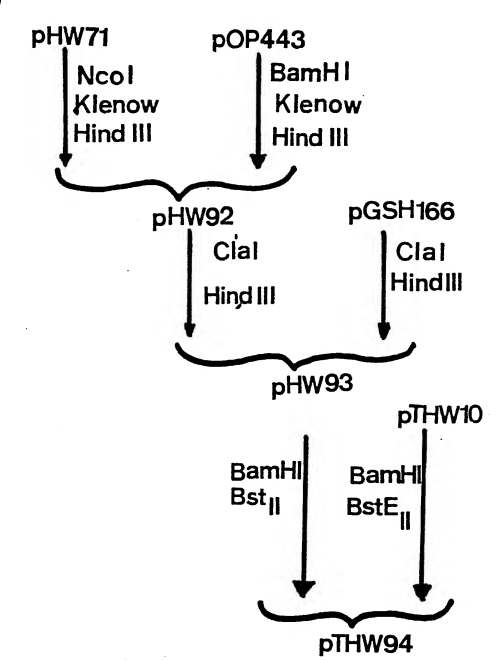
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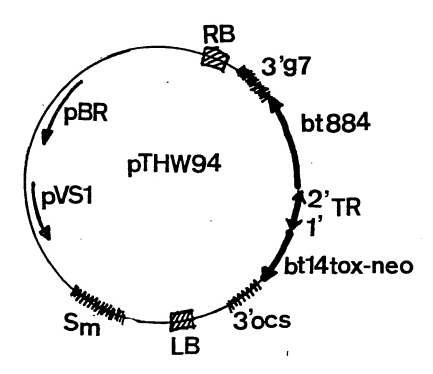
## FIGURE 16 (CONT)

d)

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# FIGURE 16 (CONT)



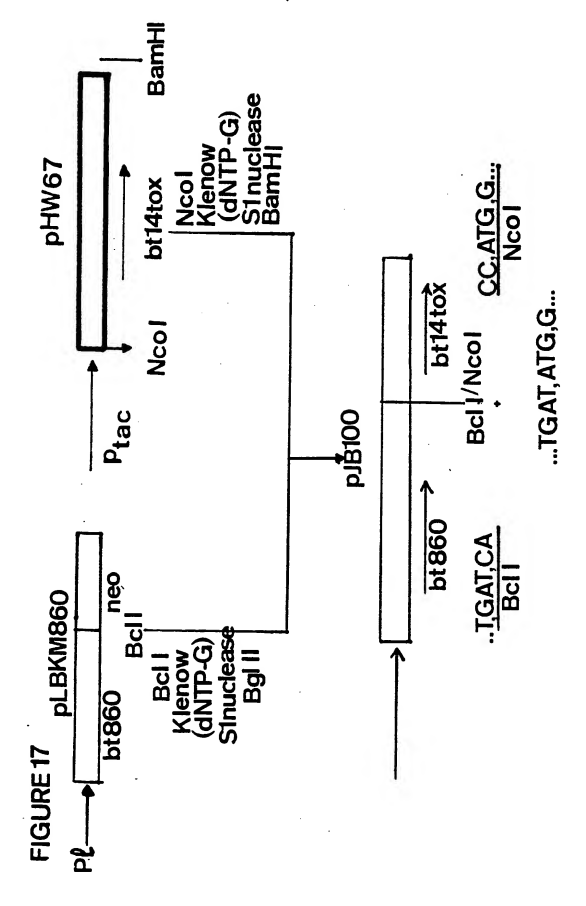
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### INTERNATIONAL SEARCH REPORT

International Application No PCT/EP 90/00905

I. CLASS	IFICATION OF SUBJECT MATTER (il several classific	ation symbols apply, indicate all) 6				
	to International Patent Classification (IPC) or to both Nation					
IPC <sup>5</sup> : C 12 N 15/32, A 01 N 63/00						
IPC :						
II. FIELDS	SEARCHED	N- C				
	Minimum Documents	assification Symbols				
Classification	on System   Ci	assincation Symbols				
IPC <sup>5</sup>	IPC <sup>5</sup> C 12 N 15/32, A 01 N 63/00					
Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched •						
III. DOCU	MENTS CONSIDERED TO BE RELEVANT		Relevant to Claim No. 13			
Category *	Citation of Document, 11 with Indication, where appro	priate, of the relevant passages 14	Relevant to Claim rec. "			
Y	EP, A, 0228838 (MYCOGEN C 15 July 1987 see the whole documen (cited in the application	it	1,13-16, 18,19			
Y	WO, A, 88/08880 (ECOGEN, INCORPORATED) 17 November 1988 see abstract; page 11, line 18 - page 13, line 12; page 25, line 9 - page 27, line 15; page 41, line 33 - page 4, line 28; claims					
A	EP, A, 0193259 (PLANT GEN N.V.) 3 September 1986 see the whole documer (cited in the application	nt	1,13-16, 18-19			
		./.				
*Special categories of cited documents: 19  *A" document defining the general state of the art which is not considered to be of particular relevance  "E" earlier document but published on or after the international filing date  "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)  "O" document referring to an oral disclosure, use, exhibition or other means  "P" 'ocument published prior to the int. document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with one or more other auch document is combined with the application but cited to under						
IV. CERTIFICATION  Date of the Actual Completion of the international Search  Date of Mailing of this International Search Report						
6th August 1990 06. 09. 90						
International Searching Authority Signature of Authorized Officer						
	EUROPEAN PATENT OFFICE	R.J. Eernisse	THE STATES			

111. DOC	III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)					
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Y	EP, A, 0192319 (MYCOGEN CORPORATION) 27 August 1986 see the whole document, in particular page 6, lines 20-24	1				
Y	EP, A, 0221024 (SANDOZ AG) 6 May 1987 see the whole document, in particular	1				
A	page 3, lines 30-34   Patent Abstracts of Japan, vol. 12, no. 391 (C-537), 18 October 1988,	1				
	& JP, A, 63137684 (SUMITOMO CHEM. CO., LTD) 9 June 1988  Patent Abstracts of Japan, vol. 13,	1				
E	no. 326 (C-620), 24 July 1989, & JP, A, 01104177 (SUMITOMO CHEM. CO., LTD) 21 April 1989					
T	Chemical Abstracts, vol. 112, no. 21, 21 May 1990, (Columbus, Ohio, US), G. Honee et al.: "A translation fusion product of two different insecticidal crystal protein genes of Bacillus thuringiensis exhibits an enlarged insecticidal spectrum", page 262, abstract 193742f, & Appl. Environ. Microbiol. 1990, 56(3), 823-5	1				

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EP 9000905 SA 37347

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